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EP 0 670 160 B1

EUROPEAN PATENT SPECIFICATION

(15)

(51) Int. Ct.6: A61K 9/46

(45) Date of publication and mention 14.07.1999 Bulletin 1999/28 of the grant of the patent:

(21) Application number: 94203112.1

(22) Date of filing: 26.10.1994

(54) Granular product or tablet containing an effervescent system and an active pharmaceutical substance, as well as a method for its preparation Ein Brausesystem und einen Arzneiwirkstoff enthaltendes granuläres Produkt bzw. Tablette sowie Verfahren zu deren Herstellung

Produit granulaire ou comprimé contenant un système effervescent et un agent actif pharmaceutique, et son procédé de préparation

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AT BE CH DE DK ES FR GB IE IT LI LU NL PT SE (84) Designated Contracting States: Designated Extension States:

(30) Priority: 01.03.1994 DE 440664: 23.03.1994 CH 87394 (43) Date of publication of application: 06.09.1995 Bulletin 1995/36

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WO-A-93/00886 US-A- 4 704 269 EP-A- 0 415 326 GB-A- 1 270 781 References cited: (26)

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Description

vescent system and a - preferably acid-sensitive - pharmaceutical substance, such as cisapride, beta-carotene, an H2 tical preparation with comparatively small amounts of effervescent components or a comparatively low acid-binding [0001] This invention relates to a granular pharmaceutical preparation or more particularly a tablet containing an efferblocker such as cimetidine or ranitidine, and/or a substance which is to be administered in an effervescent pharmaceu-

Background of the Invention

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solution, or in any event, hydrophobic particles of the drug tend to creep upward on the glass. On the other hand, in vescent tablets or effervescent instant granular products, since in contact with the acid of the effervescent system such compositions hydrolyze or decompose, i.e. they are not shelf-stable. Furthermore, whenever such a substance also affects the surface tension of water, frothing occurs which is highly undesirable for the consumption of the effervescent certain cases, the antacid side-effect of an effervescent tablet is undesirable for many drugs. Therefore an object of this invention is to provide an effervescent system which will avoid the aforesaid disadvantages and offer the possibility of administering to a patient pharmaceutical substances, inclusive of acid-sensitive substances which have hydrophobic properties or properties influencing the surface tension of water, in pleasant-to-drink effervescent solutions. It is a further object of this invention to create an effervescent tablet or an instant effervescent granular product with an acid binding capacity of less than 5 meq, in order to avoid undesired antacid effects. This is especially advantageous for all H2 olockers. Lastly, it is desired that the tablet or granular product is to dissolve rapidly in water at a temperature of about [0002] Heretofore it has been possible only with difficulty to incorporate acid-sensitive drugs in stable form into effer 15-20°C in less than about 2 minutes. 50

Summary of the Invention

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[0003] The solution to the aforesaid problems can be achieved in a surprisingly simple, cost-effective and efficient manner in accordance with this invention e.g. by first substantially coating acid particles with a corryosition contrising at least one neutral substance which causes a depression of the melling point of the acid grains al Iheir surface, and thereafter anchoring thereon at least one second coating which contains an alkali and/or alkaline earth carbonate and/or bicarbonate, and optionally a partial reaction product of the carbonate or bicarbonate with the same or a different 30

[0004] The invention is more fully discussed in detail below along with a detailed discussion and illustration of several preferred embodiments.

Detailed Description

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such as mattodextrin, dextrin and the like; especially preferred are higher alcohols, such as xylitol, mannitol and sorbitol under the influence of the only slightly alkaline effervescent grain surface due to the bicarbonate coating, are subject to nylpyrrolidone, carbohydrates, such as saccharose, pentaerythritol, glucose, and fructose (although the latter two, a Maillard reaction tending to make them yellow and therefore they are not particulary preferred herein), hydrocolloids, Neutral substances within the meaning of this invention include water soluble polymers, such as e.g. polyvi-Various embodiments of the invertion are described in the defining clauses of the dependent claims. [9002] \$

.0006] It is true that W093/00886 discloses that a foreign acid, possibly gluconic acid delta-lactone, which hydrolyzes ection for acid-sensitive active substances. It has therefore also been impossible hitherto to use the invention of to gluconic acid, can be incorporated at the surface of acid vehicle crystals, with the result that the crystal lattice is disturbed and a melting point depression is achieved. However, such a measure cannol of course provide adequate pro-WO93/00886 for acid-sensitive active substances in practice. 5

lidone described there in the Examples, foam formation problems; furthermore, some acid is always transferred from whereby the acid-sensitive active substances would not be protected sufficiently. In addition, however, those skilled in the art have for over 20 years been unable satisfactorily to solve the problem of accommodating acid-sensitive active [0007] It has also been proposed (British Patent 1,270,781) to coat acid vehicle crystals for effervescent tablets with a thin polymer layer, such as, for example, with polyvinyl-pyrrolidone, carboxymethylcellulose or the like. However, this results in an undesirable relardation of the dissolution time and, in the case of the 1 to 5% by weight of polyvinybynrothe vehicle crystal to the layer in solution when the coating is applied by means of ethanolic or aqueous solution, ow acid binding capacity and short dissolution time. An effervescent tablet is generally defined as being particularly apid when the dissolution (or complete suspending) of the tablet components takes less than 120 sec, preferably 90 substances in effervescent systems not only in a shelf-stable manner but also in relatively small tablet weights with very 20 55

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sec or less.

[0006] According to the invention, however, after (preferably only a small amount of) the neutral substance has been applied to the acid grains, alkali and/or earth alkaline carbonate and/or bicarbonate particles are anchored on the grain surface in order to prohibit an interaction between the acid and the acitive substance.

[0009] Furtilierinore, the process proposed in EP-A1-415 326 for coating acid vehicle crystals with several times the amount of sugar in order, in combination with bicarbonate, to achieve a slightly prickling effect, for a chewable tablet or locenge has not been able to solve the combination of the problems or tasks; such a system would not be sufficiently reactive to dissolve an effervescent tablet in water within a reasonable time. It was the aim of the said EP-A1 to slow down the reaction between acid and carbonate in order not to produce an undesired high effervescent effect in the

mouth.

[0010] If, as disclosed in the prior art (US-A-4 127 645), a tablet having a core of acid, bicarbonate and calcium were coated with a neutral substance, for example will soibitof solution, such a tablet would not provide reliable protection for acid-sensitive active substances contained in the core. However, if the mixture were pressed with a neutral substance (e.g. maltodextin, if necessary as a mixture will sugar, US-A-650 669; soibitol with vitamins, US-A-5 223 564, only suitable as a prickling chewable tables) to give tablets, then either both reactants would be coated together or undesirable as a prickling chewable tables) to give tablets, then either both reactants would be coated together or undesirable agglomerated granules would court, in both cases, the reaction on dissolution of the tablet would take place too slowly and the dissolution time would thus be undesirably increased, or the solution would contain undesirably large amounts of sugar. Furthermore, it is very probable that, in agglomerated granules, acid particles too would be present unprotected at the surface of the granules; however, this results in greater instability for acid-sensitive active sub-

[0011] In U.S. Patent No. 4,867,942, a method is described in which vehicle crystals of a solid, edible organic acid are covered on their surface with a pre-reacted solution serving as buffer, particularly an acid alkali and/or alkaline earth sail of a solid, edible organic acid. Thereafter, more of the acid organic and anounts of carbonate or bicarbonate are anchrored side by side on this coaling. Water which is released in the various neutralization partial reactions is removed by a final treatment with alcohol and vacuum drying. Such a process is disadvantageous, however, in that, for acid-sensitive drugs, on the acid crystal surface an additional acid simultaneously enters into a reaction with the alkali carbonate, and the reaction thus proceeds too last and consequently not sufficiently uniformly. Therefore, the product that forms from this method cannot completely prevent the reaction of an acid-sensitive drug mixed in with it, due to the acid crystals lying on the surface of the granules.

[0012] In contrast, the structure of the effervescent system according to this invention not only prevents direct contact of an acid-sensitive drug with the acid crystals thereby providing an effervescent table of granular product with substantially more shelf-stability, but it also permits the preparation of substantially smaller tablets, i.e., those with smaller amounts of effervescent components which, when dissolved, result in a buffer system. Thus, the present tablets according to the invention, in contrast to buffer systems of antacid effervescent preparations, can remain at lar less than 5 med of acid binding capacity. Aso, in terms of product preparation, a related reaction and better compressibility into tablets is obtained. With the aid of this invention, an effervescent tablet can be prepared which for the first time contains an acid-sensitive drug, such as cisapiride, or an IrB blocker such as cinnetidine, and which has an acid-binding capacity of less than 5 mey for a tablet (or granular product) weight of only 1.6 to 2.3 g.

[0013] Futtles, in accordance with an especially advantageous embodiment of this invention, after the acid crystals to have been covered with a coating of neutral substance, at least a portion of the carbonate and/or bicarbonate particles intended for a full dose can be applied to this coating, so that effervescent grains are formed from acid crystals on which a first coating of neutral substance has formed, and thereon a second coating of carbonate and/or bicarbonate, which has particles with the acid in some cases.

(0014) The invention can be particularly expediently used for products or processes as described, for example, in EPin 11-76 340, US-A-4 867 942 and WO93/00886. [0015] The application of the neutral substance, especially a sorbitol solution, for example, causes a depression of the melting point on the surface of the civic acid crystals. Thus, on the one hand, the adhesive force for the next coating containing alkeli or alkaline earth carbonates and/or bicarbonates increases, and at the same time this signifies a slower and therefore more uniform reaction of the civic acid crystal surface and better passivation, so that the acid-sensor silve drugs are less attacked by the effervescent grains. On the other hand, the melting point depression protracts the recrystalization line of the civic acid or of the circles. That have formed, which signifies better compressibility of the effervescent grainds over a longer period of time.

(0016) The amount of neutral substance applied to the acid vehicle crystals depends on the amount of solvent with which the acid can be wetled, since a maximum of 50 · 70 % by weight can be dissolved in an aqueous solution. It is therefore preferably acided in an amount of 0.05 bis 1.0, in particular 0.07 bis 0.8, % by weight, based on the acid. Addition, tone shan 0.07 have only a weak effect and those of less than 0.05 have no edited and as weak effect and those of less than 0.05 have no edited that according to the invention; the shell-stability of acid-sensitive active substances is reduced. Additions of over 0.8 generally begin to have an interfering effect, and at above 1.0 the reactivity of citic acid and of the efferedent system is considerably.

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slowed down.

[0017] However, this may be less troublesome in the case of granules since a longer dissolution time tends to be desirable there in order to allow the granules to sink on introduction into water and only thereafter to undergo a reaction for dissolution. Otherwise, however, the amounts of neutral substance which can be applied to, for example, citir acid are determined by the amount of solution with which the citir acid can be wetted, since the neutral substances are in fact applied in solution, and a 50 to max. 70% solution can be prepared. The citir acid crystals cannot be wetted with an infinitely large amount of water and hence solvent.

[0018] In certain circumstances, the neutral coaling, especially if carbonate and/or bicarbonate particles are anclored on it, can also contain small amounts of a solid, edible organic acid, and in some cases an acid different from the one of which the vehicle crystals consist - as disclosed per se in another context - but here also in order to intensity the melting point depression and/or to control the aftervescent reaction and rate of dissolution.

[0019] Each such effervescent grain is, taken by itself, actually a small effervescent "tablet", and effervesces by itself

[0020] Therefore, if desired, a short dissolving time, small quantity and low acid-binding capacity can be achieved.
[0021] Experiments thus far towards achieving a fast-acting, small effervescent tablet by the use of monosodium citis rate instead of citric acid have failed, because this greatly slows the effervescent reaction, since the monosodium citrate
reacts more slowly with sodium bicarbonate, and such tablets usually have an acid consuming capacity exceeding 5

10022] On the other hand, a very thin monosodium citrate coating in accordance with this invention, especially as a third control hayer, which can contain an additional neutral substance if desired, acts advantageously because 1 mol zo of monosodium citrate binds 1 mol of water of crystallization and thus contributes to the drying or to maintenance of dryness. Furthermore, in any case, uncovered citric acid surfaces can be covered again or more completely with bicarbonate.

[0023] Additionally, since many substances exhibit some form of taste sensation of which many are unpleasant, especially those exhibiting bitterness, it is desirable to keep the final effervescent solution, especially since it is in beverage of 50 mm, within the pH range of 3.8 to 4.6. Experience has shown that within this range paricularly bitter substances can be more effectively masked.

10024 While not obligatory, it is preferable to remove residual water from the reaction granules in the course of their preparation by a final treatment with alcohol. Alcohol may disrupt the bonding of water of crystallization, because during drying the residual moisture is removed along with the alcohol by evaporation. Small amounts of an antiboaming agent can also be added to the alcohol in order to accelerate the dissolution of the final tablet.

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10026] These bubbles burst and leave the CO₂ on the surface. Now, if a less soluble or more hydrophobic substance is present, the undissolved particles envelop the CO₂ bubbles, and by forming such a film successfully prevent rapid bubble bursting, so that the bubbles with this film on the surface collect and thus a "foam" is formed. However, the "loam" at heady forming between the effervescent grains interferes with the continued reaction, and thus with the rapid dissolution of the tablet or granules. This circumstance is combatted according to the invention by the addition of very small amounts of at least one antiloaming agent with the result that any "foam" that forms as the effervescent reaction begins immediately collapses.

[0027] The authorating agent is preferably added in an amount of 0.005 to 0.5% by weight, based on the total amount including any fillers, flavors, etc., or 0.05 - 2.0% by weight, based on active substance. Additions of less than 0.005 lave no effect relevant according to the invention; additions of more than 0.5 give rise to troublesome or unacceptable side effects.

[0028] In the case of active substances which are soluble, although not too freely soluble, as in the case of cimetidine, a percentage of simethicone of 0.1 - 0.3% by weight, based on active substance, is used, which is equivalent to the use of 0.016 - 0.028 percent (about 0.03%) based on the total tablet weight. The situation is somewhat different in the case of an insoluble hydrophobic active substance, such as cisapride (the monohydrate is used), where 1% of simethicone so is used, based on the active substance, but an amount of 0.006% results when based on the tablet weight of 1.6 g. It is evident that the cisapride, as a slightly soluble, hydrophobic active substance, requires a farger amount of antifloaming agent for suppressing the foam, but the required filters and the effervescent base result in a substantially smaller amount of simethicone being used per tablet, so that the ratios are inverted.

[0029] In the case of the soluble active substances, such as cimetidine and ranifidine, the sinethicone is required in smaller amounts, in order to suppress the smaller tendency to loaming in the local reaction on dissolution of the effervescent tablet, whereas in the case of cisapride - as already mentioned - the tendency to loam is substantially greater and the principle is therefore also slightly different.

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0030] If larger amounts are used, film formation of simethicone occurs at the surface after dissolution of the efferves-

stance collect and remain hanging and thus result in unattractive dissolution behavior, this film then additionally having cent tablet, by virtue of the fact that - especially in the case of insoluble active substances - particles of the active subthe tendency to form a ring on the glass wall.

- 10031] In some cases, however, very small amounts of a tenside, for example, docusate sodium, are also added. Due to their wettable nature, such drug particles dissolve more quickly and no longer adhere to the foam bubbles. The proportion of such substances must be determined very precisely to achieve the desired dissolving characteristics.
- .0032] Although in some cases the antifoaming agent can be applied to the effervescent system and/or to the drug, his is not preferred according to the invention. In the first case, it might cause undesirable slowing of the dissolution ment of the desired effect are used. In the second case, only those drugs are involved which, when the antifoaming agent is drawn onto them from a solution in a solvent (e.g., methyl ethyl ketone and acetone) at 40°C, do not lose any of their solubility or stability. Additionally, in the course of production with the use of finely powdered drugs the addition of antifoaming agents may lead to poor distribution because of drug particles attaching themselves to the antifoaming and reaction of the effervescent components unless very slight amounts of antifoaming agent sufficient for the achieveagent droplets.
- (0033) It is therefore preferred, in accordance with this invention, that first the formation of a typical granular product from antifoaming agents and a neutral substance is undertaken, which product is thereafter mixed with the effervescent system and the drug, plus additional adjuvants if desired (e.g., perfumes, sweeteners and the like) and the mixture then compressed into tablet form.
- [0034] The moisture released in the preparation of the effervescent system by the neutralization reaction, and not entirely removed by heating and/or vaccuum treatment, as well as moisture picked up from the air during storage, can best be bound by the addition of a moisture-binding agent, especially anhydrous sodium carbonate (which can absorb 10 mols of water per mot) or sodium sulfate. The agent can be bound either by applying it to one or more of the coatings applied to the vehicle crystals, or by adding it to the total mixture. This improves shelf life because the reaction of the acid-sensitive active agent with the acid is further suppressed or completely prevented by the reduction of moisture. However, excessive amounts of such moisture-binding agent, for example sodium carbonate, are not desirable as it may retard the effervescent reaction. 20
 - grains, since it is preferable to operate with only small quantities effective to merely dry the residual moisture, or to Sodium carbonate as a drying agent, therefore, should not be used for completely covering the effervescent retard the reaction during manufacture, and to avoid undesirably lengthening the dissolving time of the tablet. Therefore, [0035]
- the final addition of sodium carbonate should not be used for complete coverage (or a tablet coating), due to both the quantity and the grain size (approx. 0.1 - 0.05 mm), and it is therefore not suitable for producing a continuous coating on the bicarbonate already present. However, it can be partially hooked onto the effervescent grains. It is also possible, however, not to add the sodium carbonate until after the drying operation. 8
- In principle, the percentage amount of sodium carbonate per tablet depends on several factors, such as, for example, the amount of effervescent base used, the amount and type of the fillers used, the presence of other carbonates, such as, for example, calcium carbonate, etc. [0036]33
- and 10, in particular 4 6, % by weight (based on the total amount, including any fillers, flavors, etc.). Additions of less than 4 have only a weak effect, and with those of less than 1, the drying effect and increase in stability is too small, they have no effect relevant according to the invention. Additions of over 6 generally begin to have a troublesome effect significantly lengthened, since sodium carbonate first absorbs water (up to 10 molecules of water of crystallization) on [0037] The moisture-binding agent, in particular sodium carbonate, is preferably added in an amount of between 1 because sodium carbonate dissolves more slowly and reacts more poorly; above 10% the dissolution time is already dissolution of the effervescent tablet, i.e. is calcined and only then reacted with the citric acid. 5
 - [0038] Here it is to be emphasized that 1 mol of water of crystallization can be bound per mol by sodium citrate atone developing in or on the sorbitol layer, and in spite of any residual moisture present the sorbitol layer prevents or hinders any acid harm to the drug. 5
- duced, even with the difficult substances referred to, which at a tablet weight of, e.g., 1.6 g,will attain a dissolving time If all of the prescribed steps are followed in accordance with the invention, effervescent tablets can be proof less than 100 seconds. It is also to be noted that especially cimetidine, due to its hydrophobic character, further lengthens the dissolving time in comparison with other drugs, under otherwise equal conditions. [0039]
- [0040] Granulation with sorbitol solution permits rapid dissolution without the incorporation of an extraneous acid that is otherwise necessary, for example, according to WO93/00886.

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- [0041] Furthermore, during the preparation of the effervescent systems of this invention, and in any case of the tablets face of individual crystals or granules, which thus constitute a local mechanism, while also during dissolution the abovethemselves, the steps taken according to the invention will enable the control of reactions which take place at the surdescribed desired advantages will be achieved throughout. £
- [0042] The system is also extraordinarily well suited for the processing of substances which are both acid-sensitive and sparingly soluble in water. Such substances, such as cisapride for example, exhibit very unpleasant behavior in

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suspension, since, as mentioned above, they tend to froth together with the effervescent system, adhere to a glass wall, form unpleasant rings and tend to agglomerate on the surface of the drink.

- All the aforesaid problems can be effectively combatted by preparing seperate granules. For this purpose in yet another embodiment of this invention, there is provided a vehicle which can consist of an Aerosil and/or a neutral substance, on which the drug is applied preferably with the surface of its grains partially dissolved, and/or with birkfing
- 19044] The amount of the suspended substance is limited to at most 8, preferably at most 4.5, % by weight (based on the total mixture), for example for cisapride, since larger amounts would result in increased sinking of the granule paragents and/or tensides if desired, and dried, or is bound to the vehicle surface by means of binders.
 - ticles after dissolution of the tablet. On the other hand, the amount of binder used is likewise limited to 1% by weight, since it otherwise leads to undesirable agglomerated granules of active substance, susperxted substance and binder, which dissolve only with difficulty and then sink to the bottom, i.e. prevent the desired suspension. 9
 - [0045] The invention will now be more fully described and understood with reference to the following examples of prelerred embodiments.
 - [0046] _Alternatively, the drug can also be dissolved in the methyl ethyl ketone or in acetone and coaled onto mannitol Aerosii^(F) and sodium bicarbonate. 5

Example 1: Preparation of effervescent tablets containing 200 mg of cimetidine

a) Preparation of the effervescent system 8

- face on which up to about 30% of bicarbonate can be anchored) or tartaric acid are aspirated into a preheated vacuum tank and heated to approx. 60°C with stirring. Next, 0.85 parts by weight of a solution 1, which has been formed from 36 parts by weight each of water and sorbitol, 21 parts by weight of citric acid and 7 parts by weight of sodium bicarbonate, is aspirated and distributed on the citric acid by mixing. Thereafter, 52.5 parts by weight of sodium bicarbonate and 4.4 parts by weight of aspartame are added to this mixture, which is then stirred and dried by a vacuum of up to 200 mbar, after which 1.9 parts by weight of sodium carbonate are aspirated and dishibuted in the mixture by stining. erable for improving build-up to effervescent grains on the vehicle crystal as the powder particles provide a rough sur-102 parts by weight of coarse citric acid and 25 parts by weight of finely powdered citric acid (the latter is prefand the mixture is then dried by a vacuum of up to 15 mbar.
- [0048] Next, a further 0.6 parts by weight of said solution are aspirated and distributed in the mixture by mixing. The resultant effervescent grains are dried in a vacuum of up to 20 mbar with stiming. If necessary, 0.25 parts by weight of 96% eithand are also employed to dry the mixture, and aspirated. Then, again 9.3 parts by weight of sodium carbonate are bound onto the effervescent grain surface. After another final drying, the product is removed through a sieve. 30

b) Preparation of the granulated antifoaming agent 35

In a vacuum mixing tank with a jacket temperature of 80°C, 7.7 parts by weight of sorbitol powder are added and heated to 50°C Then, 0.2 parts by weight of simethicone in a 30% butanone/acelone mixture (5:3) are aspirated in, stirred by vibrational mixing and dried under full vacuum down to 15 mbar at a temperature of at least 45°C.

c) Preparation of the total mixture

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for 10 minutes at 6 rpm with 178.4 parts by weight of the effervescent system prepared in a). Then 7 parts by weight of the antifoaming agent granules prepared in b) and screened through a 0.6 mm sieve, and 4.5 parts by weight of lemon [0050] In a mixer, 20 parts by weight of cimetidine, with 21.1 parts by weight of sorbitol powder if desired, are mixed flavoring, are added, mixed for another 5 minutes at 6 rpm. The final mixture is pressed into tablets which weigh 2.3 g. contain 200 mg of cimetidine, and have a hardness of 6-8 kp. 45

Example 2: Preparation of effervescent tablets containing 200 mg of cimetidine, and citric and matic acid in the effervescent grains: 20

added to the mixture and dried by stirring, in a vacuum of up to 200 mbar. Next, 1.9 parts by weight of sodium carbonate are aspirated in and distributed in the mixture by stirring, and then vacuum drying is performed down to 15 mbar. Finally, 102 parts by weight of coarse citric acid, 25 parts by weight of powdered citric acid and 1.1 parts by weight of malic acid are heated to 60°C with stirring in a preheated vacuum tank. A solution consisting of 0.4 parts by weight of water, 0.22 parts by weight of sorbitol and 0.22 parts by weight of malic acid is then aspirated in and distributed onto the citric acid by mixing. 52.5 parts by weight of sodium bicarbonate and 4.4 parts by weight of aspartame are next a final drying can be performed with ethanol, and then 9.3 parts by weight of sodium carbonate are added to the mix-22

ture. The rest of the procedure is similar to Example 1.

Example 3: Effervescent tablets containing 400 mg of cimetidine, and mannitol as a neutral substance

6 [0052] 49 parts by weight of olitic acid are aspirated into a preheated vacuum lank and heated with stinring to 60°C. Then, 0.45 parts by weight of a solution 1, which has been prepared from 0.25 parts by weight of water and 0.20 parts by weight of naminol, is aspirated in and distributed on the citie acid by mixing, whereupon 14.7 parts by weight of social parts by weight of aspartame are then acided Reaction is started with stirring and then dry ing is performed with a vacuum to 15 mbar. Then 0.5 parts by weight of aspartame are then acided Reaction is started with stirring and then dry ing is performed with a vacuum to 15 mbar. Then 0.5 parts by weight of a solution 2, which has been prepared from solution 1 by the aciditor of 0.16 parts by weight of monosocium citrate, is aspirated into the instrure and distributed by mixing. The effervescent grains obtained therefrom are then dired by vacuum and stirring a 20 mbar, and finally 2.8 parts by weight of social carbonate are added. To this mixture are then acid from a parts by weight of social carbonate are added. To this mixture are then acid from a parts by weight of mannitol, 8 parts by weight of antificanting agent granules prepared according to Example 1 b), until distribution is uniform.

Example 4: Effervescent tablets containing 300 mg of cimetidine, as well as mallodextrin as a neutral substance

20 [0053] Similarly to Example 3, for a 300 mg cimetidine effervescent tablet, a 50% solution of maltodaxtrin is selected, which is then used in the same amount as in the case of the 400 milligram form.

[0084] In all of the examples in which the tablets contain 100 to 400 mg of cimetidine, the tablet weight can be 2.3 g. The tablets have a dissolving time of preferably 60 to 150 seconds and a buffering capacity below 5 meq, measured according to USP XXII, by back-titration (with 0.5 N NaOH) of an effervescent tablet dissolved in 70 ml of water and with 25 30 ml of 1.0 N HCl added.

[0055] The figures given in the following table 1 are the percentages of individual ingredients in the particular total mixture of the illustrated preferred embodiments, which therefore are in the following preferred ranges:

Table 1

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Cimetidine	2 - 30%	(corresponds tablet containin cim	(corresponds to an effervescent tablet containing 50 to 600 mg of cimetidine)
Citric acid	30 - 60%	sorbitol	5-20%
Sodium bicarbonate	10 - 30%	mannitol	2-10%
Sodium carbonate	2 - 10%	simethicone	0.005-0.5%
Aspartame	1-4%	flavoring	0.1-3%

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[0056] A preferred percentage composition of cimetidine effervescent tablets or bags of granules containing 100, 200, 300 and 400 mg of cimetidine, with a total weight of 2.31 grams, is summarized below in table 2:

Table 2

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	•		E	Ē.
Cimetidine 4.3	4.30	8.70	13	17.3
Citric acid 50		20	48.2	48.2
Sodium citrate 0.0	0.04	0.04	0.04	0.04
Aspartame 1.7	1.74	1.64	2.54	3.24
Sorbitol 12.5	2	12.5	12.8	8.00
Sodium bicarbonate 20.7	_	20.7	14.7	14.7
Sodium carbonate 4.4	4	4.4	3.5	3.3

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Table 2 (continued)

	100 mg	200 mg	300 rng	400 mg
Manntiol	4.3		4.3	4.3
HMA Lemon flavoring	2.0	2.0	6.0	6.0
Simethicane	0.02	0.02	0.02	0.02

Example 5: Cisapride effervescent tablets

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a) Preparation of the effervescent grains

[0057] 655 parts by weight of crystalline citric acid, 70 parts by weight of citric acid powder and 8 parts by weight of scolum saccharin socium are heated while mixing to 60°C. Then 28 parts by weight of socium are heated while mixing to 60°C. Then 28 parts by weight of citric acid and 1.6 parts by weight of citric acid and 1.6 parts by weight of citric acid and 1.6 parts by weight of weight of sociation to an order of the mixing to 60°C. Then 28 parts by weight of sociation bicarbonate as well as 2 parts by weight of sociation bicarbonate as well as 2 parts by weight of sociation bicarbonate as well as 2 parts by weight of sociation carbonate are added, whereupon the mixinc also well as 10°C. That is by weight of sociation carbonate are added whereupon the mixinc as well as 10°C. That is a 10°C. That is the weight of sociation carbonate are

20 b) Preparation of the granulated drug

[0058] Insoluble and hydrophobic disapride is attached to a suspending substance by means of a binder and a small annount of a tenside as follows: A solution of 10 parts by weight of disapride, 2 parts by weight of polyvinylpyrrolidone and 0.8 part by weight of docusate sodium in 1 part by weight of than and 40 parts by weight of acetone is applied as to 10 parts by weight of Aerosif^(R), uniformly distributed and then dried while stirring. The granules are sieved to 0.1 -

c) Preparation of the end mixture

20 (19059) To 1152 parts by weight of effervescent grains are added 50 parts by weight of maltodextrin, 100 parts by weight of lactose, 184 parts by weight of mannitol, 40 parts by weight of lactose, 184 parts by weight of mannitol, 40 parts by weight of lactose, 184 parts by weight of anti-foraming granules (0.2 parts by weight of simethicone coated onto 50 parts by weight of mannitol), as well as the granulated drug prepared in b), mixing is carried out for 15 minutes for uniform distribution and the nixture is then pressed to form tables of 1.6 g, which have an acid-binding capacity of only 2 meq. Cisapride effervescent tablets having such a low acid-binding capacity are unknown to date.

Example 6: Beta-carotene effervescent tablets

10060] With this extremely acid- and oxidation-sensitive aubstance, attention must be paid to an especially good covering of the acid. The surface and the contact zone on the beta-cardene must be kept alkaline. Therefore the effervescent grains are covered at least in part with catcium carbonate, thus insuring an alkaline surface. This, however, does result in a slightly longer dissolving time, which in this case is desirable, because the beta-cardene needs time to suspend while the tablet is dissolving. Large amounts of societic, as in US-A-5.22.264 mentioned at the outset, are by no means suitable for a beta-cardene effervescent tablet which is interded to be dissolved or suspended in water.

a) Preparation of the effervescent grains

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[0061] 1315 parts by weight of citric acid, 7 parts by weight of sodium saccharin and 45 parts by weight of sodium cyclamate are heated in a vacuum tank to \$0°C. Then 168 parts by weight of a solution prepared from 3.6 parts by weight of calcium carbonate, 19 parts by weight of solution prepared from 3.6 parts by weight of calcium carbonate, 19 parts by weight of citric acid, 12 parts by weight of sorbitol, and 45 parts by weight of water are stimed in and distributed onto the olicin acid by mixing. Next, 400 parts by weight of calcium carbonate and 190 parts by weight of calcium carbonate and 190 parts by weight of the above-mentioned solution, and after distributing and mixing, 403 parts by weight of solution are acided and the mixture heated with stirring to 60°C. Then follows the second gianulation with 44 parts by weight of the above-mentioned solution, and after distributing and mixing, 403 parts by weight of sodium bicarbonate and also, before drying, 52 parts by weight of sodium carbonate. The mixture is then secunn-dried to 15 mbar with slow mixing.

b) Preparation of the end mixture

encapsulated beta-carotene suspendable in water and corresponding to 2 to 15 parts by weight of 100% beta-carotene, 130 parts by weight of sorbitol and 540 parts by weight of mannitol and 50 parts by weight of flavoring, an spunding to 10 to 75 parts by weight of 100% tocopheryl acetate), plus still other vitamins if desired, are mixed with 2415 parts by weight of the effervescent grains prepared according to a). The product has a tablet weight of 3.3 g and plus, it desired, 50 to 250 parts by weight of vitamin C and/or a solid tocopheryl acetate suspendable in water (correits dissolving time is 60 to 90 seconds.

Example 7: Ranitidine effervescent tablets

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a) Preparation of the effervescent grains

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840 parts by weight of crystalline citric acid, 210 parts by weight of citric acid powder, 45 parts by weight of sodium cyclamate, and 4 parts by weight of sodium saccharin are heated in a vacuum mixing tank at 60°C. Then a solurated in and distributed by stirring. 500 parts by weight of sodium bicarbonate are next added and allowed to react, and ion consisting of 6 parts by weight of water, 1 part by weight of sodium citrate, and 3 parts by weight of sorbital is aspithereatter 370 parts by weight of monosodium citrate are added, which are also allowed to react. Lastly, 100 parts by weight of sodium carbonate are added and the granules are dried with slow stirring up to 15 mbar. [0063]

b) Preparation of the end mixture

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[0064] To the effervescent grains thus prepared, 167 parts by weight of ranitidine hydrochloride, 125 parts by weight of mannitol plus 100.4 parts by weight of a granulated antitoaming agent (consisting of 100 parts by weight of mannitol and 0.4 parts by weight of simethicone) and the flavoring agent are added. This mixture is mixed for 15 minutes for unilorm distribution, and then pressed to tablets of 2.5 g. The tablets have a dissolving time of 60 to 80 seconds and an acid-binding capacity of about 2 meg and contain (in percent by weight) 6.8 ramitdine hydrochloride, 42.0 citric acid, 14.8 monosodium citrate, 20.0 sodium bicarbonate, 4.0 sodium carbonate, 2.0 sweeteners, 5.0 mannitol, 0.1 sorbitol, 4.0 granulated antifoaming agent (containing 0.016 dienthylpolysiloxane) and 1.2 flavoring.

Example 8:

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voting, until uniform distribution is obtained. The mixture is then pressed to tablets weighing 143 g and having a dis-solving line of 65-70 sec, a hardness of 8 kg, and an acid-binding capacity of about 1.5 meq. The product contains no monosodium citrate. Ramitdine effervescent tablets having such a low acid-binding capacity have not been disclosed 545 parts by weight of crystalline citric acid and 133 parts by weight of powdered citric or tartaric acid are mixed while heating to 60°C. Then, as the first coating, a solution which consists of 6 parts by weight of water and 4 parts by weight of sorbitol is distributed on the surface by stirring. Next, 222 parts by weight of sodium bicarbonate are made to react on the surface of the citric acid, and finally 80 parts by weight of sodium bicarbonate are added. The product is dried with slow stirring. The granules are screened to 1.5 mm, and then mixed for 10 minutes at 10 rpm with 167 parts by weight of raviidine hydrochloride, 100 parts by weight of anti-loaming granules (containing 0.4 parts by weight of simethicone and 100 parts by weight of factose), plus 54 parts by weight of sweetener and 40 parts by weight of fla-[0065] to date 3

Example 9: £

[0066] 38.2% of citric acid is healed with 0.26% of sodium saccharin to 60°C, then two-thirds of a solution which consists of, with respect to the final mixture, 0.6% water, 0.18% sorbitol, and 0.12% sodium citrate are applied. The solution ting with the third one-third of the solution; then 12.9% monosodium citrate and, finally, 5.2% sodium carbonate are 50°C, to 15 ritbar. The basic effervescent granular product is screened to 1.5 mm and mixed with 11.0% of ranitidine lydrochloride, 6.5% of mannitol, 6.5% of anti-foaming granules plus 0.2% of flavoring, and pressed to tablets of 1.55 g. is distributed for 5 mirutes by mixing at 10 rpm. Then 16.2% of sodium bicarbonate and 2.9% of aspartame are added and anchored on the surface of the citric acid by reaction on the neutral substance coating. Then follows a second wetadded. The effervescent grains are dried while mixing them slowly by applying a vacuum, at a temperature of at least which have a dissolving time of 50 sec at a hardness of 7.3 kp and an acid-binding capacity of less than 2 meq. 95 55

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Example 10: Vehicle crystal grains coated only with a neutral substance

Since cisapride, for example, in comparison to ramifdine, is not as highly sensitive to acid, it is nevertheless also possible by the procedure to be described below to achieve protection against the acid, all the more so since the drug is embedded in granutes.

a) Preparation of the acid crystals coated with a neutral substance

Then a solution of 4 parts by weight of sorbitol in 4 parts by weight of water is applied and distributed onto the surface 593 parts by weight of crystalline citric acid plus 70 parts by weight of citric acid powder are heated to 60°C. of the citric acid by mixing. Finally the citric acid thus coated is vacuum dried at 50 to 60°C. [8900] 10

tain a second alkali or alkali earth carbonate coating, it is possible to protect cisapride, for example, against attack by 10069] In the case of both the form of effervescent product presented here and that of effervescent grains which conthe citric acid in the drug granules by the addition of sodium bicarbonate.

b) Preparation of the drug granules

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K30, 1 part by weight of propylene glycol and 0.8 parts by weight of docusate sodium are added and distributed for 5 160 parts by weight of mannitol, 10 parts by weight of cisapride, 5 parts by weight of aerosil and 10 parts by weight of sodium bicarbonate are heated with mixing to 60°C. Then half of a solution of 27 parts by weight of methyl minutes for the purpose of uniform wetting. The mixture is dried to 0.8 bar, the second part of this solution is aspirated, ethyl ketone (or 45 parts by weight of acetone), 2 parts by weight of alcohol, 2 parts by weight of poly(vinyl pyrrolidone) and again distributed by stirring for 5-10 minutes, and finally vacuum dried. [0000] 2

neutral substance, the remaining carbonates and bicarbonates, as well as the other tablet ingredients, and pressed to [0071] The active agent granules are then screened to 0.3 mm and already have an enhanced protection against acid attack simply due to the sodium bicarbonate they contain. They can then be mixed with the acid crystals coated with give tablets. £

c) Preparation of the end mixture

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[0072] The citric acid dried and coated according to a) is then mixed with the drug granules prepared according to b), 50 parts by weight of sweetener, 80 parts by weight of sodium carbonate, 430 parts by weight of sodium bicarbonate, and 50 parts by weight of maltodextrin, 100 parts by weight of lactose, 150 parts by weight of mannitol, 50 parts by weight of an antitoaming granulate, and 20 parts by weight of flavoring, and then pressed to tablels of about 1.6 g, which have a dissolving time of 60 to 70 seconds at a hardness of 7 kp. 33

Example 11: Cisapride effervescent tablets

a) Preparation of the effervescent granules:

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0.45 part by weight of cliric acid and 1.2 parts by weight of water. 12 parts by weight of matic acid are then aspirated in and uniformly anchored on the sorbitol layer formed on the cliric acid crystals. Finally, 205 parts by weight of socilum bicarbonate and 1.2 parts by weight of aspartame are aspirated in and once again uniformly distributed. Finally, the 40 parts by weight of powder, together with 5 parts by weight of saccharin sodium, is uniformly wet at 60°C with 2.2 Citric acid, consisting of an amount of 300 parts by weight of granules, 80 parts by weight of fine granules and parts by weight of a solution which contains 0.4 part by weight of solbitol, 0.15 part by weight of sodium bicarbonate, material is covered with 46 parts by weight of sodium carbonate, vacuum-dried and discharged through a 1.2 mm sieve. [0073] \$

b) Preparation of the active ingredient granules:

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[0074] 12 parts by weight of polyvinylpyrrolidone are dissolved in 12 parts by weight of ethanol; 6 parts by weight of propylene glycol and 6 parts by weight of docusate sodium are then added and the mixture is diluted with 165 parts by weight of ethyl methyl ketone. Half of this solution is distributed over a mixture of 960 parts by weight of mannitol, 30 parts by weight of Aerosil^(R), 60 parts by weight of sodium bicarbonate and 61 parts by weight of cisapride, which is heated to 60°C. Partial drying is then carried out in vacuo, and further wetting is effected with the second half of the solution, followed by complete drying and dicharge through a 0.3 mm sieve. 55

The end mixture is prepared analogously to Example 5.

Claims

- 1. A granular effervescent product suitable for preparing an aqueous solution or suspension of one or more pharmaceulically active substances for oral administration, being capable of being pressed into tablets, and/or said product in tablet form, comprising effervescent grains obtained from carrier crystals of at least one solid, edible organic acid which are substantially covered by at least one coating containing at least one water-soluble neutral substance, wherein said neutral substance is effective for depressing the melting point of the acid crystals on their surface, and at least one substance selected from the group consisting of alkali carbonate, alkali bicarbonate, alkaline earth sair bonate, alkaline earth sair one solid edible organic acid is applied onto said coating.
- The granular product or tablet according to claim 1, wherein the neutral substance is selected from the group consisting of a water-soluble polymer, a higher alcohol, a carbohydrate and a hydroxolloid, and which neutral substance is present in an amount of from about 0.05 to about 1.0 % by weight, preferably from about 0.07 to about 0.8 % by weight.

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- The granular product or tablet according to daim 1 or 2, wherein a moisture-binding agent is anchored on said
 effervescent grains, which moisture-binding agent preferably is selected from the group consisting of anhydrous
 sodium carbonate and sodium sulfate and preferably is applied in an amount of from about 4 to about 10 % by
 weight with respect to the total mixture.
- 4. The granular product or tablet according to any one of the preceding claims, wherein on the effervescent grains at least one additional coating is applied, comprising a substance selected from the group consisting of alkali salts and/or alkaline earth salts of at least one solid, edible, organic acid as buffer and, optionally, comprising an additional neutral substance, and wherein preferably at least one of the coatings contains an antifoaming agent.
- The granular effervescent product or tablet according to any one of the preceding claims, wherein the granular product, or said granular product compressed in tablet form, further comprises at least one antifoaming agent' present in a granular product of its own.
- The granular product or tablet according to daim 4 or 5, wherein the antifoaming agent is selected from the group consisting of dimethicone and simethicone and is applied in an amount of from about 0.005 to about 0.5 % by weight with respect to the total mixture or from about 0.05 to about 2.0 % by weight with respect to the pharmaceutically active substance.
- The granular product or tablet according to any one of the preceding claims, wherein it has an acid-binding capacity of less than 5, preferably less than 3 meq, measured according to USP XXII.
- The granular product or tablet according to any one of the preceding claims, wherein, at a total weight of no more
 to than 2.5, preferably no more than 2.0 grams, in water at room temperature, it has a dissolving time of less than
 180, preferably less than 120 seconds.
- 9. The granular product or tablet according to any one of the preceding dainns, comprising a pharmaceutically active substance which is hydrophobic and wherein the hydrophobic substance is present in granules separate from the effer vescent components, in which granules the hydrophobic substance is coated or anchored onto at least one substance selected from the group consisting of suspending agents preferably selected from the group consisting of Aerositing of Aerositing of mannitol and sorbito.
- 10. The granular product or tablet according to claim 9, wherein the granules also contain at least one component selected from the group consisting of binders preferably polyvinylpyrrolidone (PVP) -, small amounts of a tensifier preferably selected from the group consisting of dioctyl sodium sulfosuccinate and sodium lauryl sulfate -, alkali and/or alkaline earth carbonate and/or bicarbonate.
- 11. The granular product or tablet according to any one of the preceding claims, wherein it contains, with respect to the total nixture, about 2 to about 30 % by weight of clmetidine; about 30 to about 60 % by weight of a solid, edible organic acid; about 12 to about 40 % by weight of at least one alkali or alkaline earth carbonate or bicarbonate (of which about 2 to about 10 % by weight is sodium carbonate as moisture-binding agent); about 1 to about 4 % by

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weight of a sweetener; about 0.01 to about 30 % by weight of a neutral substance (of which about 0.01 to about 0.05 % by weight is for the neutral substance ocating), preferably about 3 to about 20 % by weight of sorbitol and about 2 to about 10 % by weight of mamitol; about 0.005 to about 0.5 % by weight of mamitol; about 0.005 to about 0.5 % by weight of an antitioarning agent, and about 0.1 to about 3 % by weight of flavoring agent, and

- 12. The granular product or tablet according to any one of daims 1 to 10, wherein it contains, with respect to the total mixture, the following components: about 0.4 to about 4.5 % by weight of clsapride; about 0.4 to about 0.5 % by weight of suspending agent; about 0.1 to about 0.8 % by weight of therefore preferably diodyl sodium sulfosuccinate; about 30 to about 0.53 % by weight of tenside, preferably diodyl sodium sulfosuccinate; about 30 to about 55 % by weight of tenside, preferably diodyl sodium sulfosuccinate; about 30 to about 55 % by weight of tenside or alkaline earth carbonate or bicarbonate (of which about 12 to about 10 % by weight of alterest on alkaline such about 0.3 to about 2.5 % by weight of sweetener; about 0.02 to about 0.3 to about 0.2 % by weight of neutral substance (of which about 0.02 to about 0.1 % by weight is for the neutral substance coating), preferably selected from the group consisting of mallodextrin, lactose and mannitiot; about 0.05 to about 0.2 to about 5.8 % by weight of antifoaming agent, preferably selected from the group consisting of dimethicone and simethicone; and about 0.2 to about 5.8 % by weight of lavouring.
- The granular product or tablet according to any one of claims 1 to 10, wherein it contains, with respect to the total mixture, the following components:
- about 0.1 to about 0.5 % by weight of beta-carotene (100%);

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- about 0 to about 2 % by weight of tocopheryl acetate (100%);
- about 35 to about 70 % by weight of solid, edible organic acid, preferably about 0 to about 10 % by weight of
 ascorbic acid, about 35 to about 55 % by weight of citric acid, and about 0 to about 5 % by weight of malic acid, and about 0 to about 5 % by weight of malic acid.
- about to about 38 % by weight of at least one alkali or alkaline earth carbonate or bicarbonate, preferrably about 15 % by weight of calcium carbonate and about 20 % by weight of calcium carbonate and about 5 to about 20 % by weight of sodium bicarbonate.
- about 1 to about 4 % by weight of sweetener;
- about 0.1 to about 35.0 % by weight of neutral substance (of which about 0.1 to about 0.5 % by weight is for the neutral substance coaling), preferably about 1 to about 10 % by weight of sorbitol and about 5 to about 25 % by weight of mannitol: and

- about 0.3 to about 3 % by weight of flavouring.
- 14. The granular product or tablet according to any one of claims 1 to 10, wherein it contains, with respect to the total mixture, the following components: about 3 to about 14 % by weight of rantitidine hydrochloride (75 300 mg per dose); about 30 to about 50 % by weight of clitic acti; about 0 to about 20 % by weight of morrosodium citrale; about 10 to about 30 % by weight of sodium bicarbonate, about 2 to about 10 % by weight of sodium carbonate; about 10 about 3 % by weight of sodium carbonate; about 10 about 3 % by weight of swelener; about 0.5 to about 0.2 % by weight of neutral substance for the first coaling as well as about 0 to about 15 % by weight of additional neutral substances of the first of antifloaming granules, and about 0.1 to about 4 % by weight of flavoring.
- 15. An effervescent tablet containing at least one pharmaceutically active substance and an effervescent system comprising at least one solid, edible, organic acid, at least one alkali metal carbonate or bicarbonate as a gas-forming component and at least one alkali metal salt of the acid, at least two layers being applied to carrier crystals constituted of the at least one acid, wherein the first layer contains at least one obtain, solid, edible, organic acid or the alkali metal salt of this other acid, or both, whereas the sectord layer contains at least one alkali metal salt of said at least one acid, and wherein the first layer additionally contains a neutral substance selected from the group consisting of a water-soluble polymer, a higher alcohol, a carbohydrate and a hydrocolloid.
- 50 16. A granular product or tablet with an effervescent system according to any one of claims 1 15 and cisapride as the pharmaceutically active substance, wherein, at a total weight of less than 2 grants, preferably less than about 1.6 grants, it has an acid-binding capacity of less than 5 meq, preferably less than 3 meq.
- 17. A granular product or tablet with an effervescent system according to any one of claims 1 15 and crimetidine as the pharmaceutically active substance, wherein, at a total weight of less than 2.5 grams, preferably less than about 2.0 grams, it has an acid-binding capacity of less than 5 meq, preferably less than 3 meq.
- 18. A granular product or tablet with an effervescent system according to any one of claims 1 15 and rauffidline as

the pharmaceutically active substance, wherein, at a total weight of less than 2.6 grams, preferably less than 2.0 g, it has an acid-binding capacity of less than 3 meq, preferably less than 2 meq.

- 19. A method for the preparation of a granular product or of a tablet according to any one of the preceding claims, wherein crystals of at least one solid, edible organic acid are wetted with an aqueous solution of a neutral substance, and then, before complete drying, an alkali and/or alkaline earth carbonate and/or bloarbonate in powder form is unitamity distributed and anchored on the moist surface layer by mixing, whereupon the effer-vescent grains thus prepared are dried and mixed with a pharmaceutically acide substance preferably with an acid-sensitive one especially one that is selected from the group consisting of H2-blockers, climetidine, rantifidine clispride and beta-carolene and pharmaceutically acceptable adjuvants, and optionally compressed into tablets.
- 20. The method according to claim 19, wherein, on the effervescent grains, at least one additional coating is applied by welling the grains with the solution of a buffer substance, preferably one that is selected from the group consisting of alkalic carbonate, alkalic bicarbonate, alkalic bicarbonate, alkalic bicarbonate, alkalic and earth said one soid exible organic acid and alkaline earth said of a least one soid exible organic acid and alkaline earth said of a least one soid exible organic acid.

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- The method according to claim 19 or 20, wherein the solution further comprises a neutral substance selected from the group consisting of a water-soluble polymer, a higher alcohol, a carbohydrate and a hydrocolloid.
- 22. The meltod according to any one of claims 19 to 21, wherein, in addition to the drug, the effervescent grains are also mixed with a granular product which has been made by applying an antitoaming agent in an appropriate solvent to the surface of neutral substance particles, and drying the solvent.
- 23. The method according to any one of claims 19 to 22, wherein the dried effervescent grains are wetted with ethanol, which preferably contains an antifoaming agent dissolved, and are dried again, by evaporating the ethanol, to remove residual moisture.
- 24. The method according to any one of claims 19 to 23, wherein the pharmaceutically active substance, before admixing it to the effervescent system, is together with a binding agent and/or a tenside applied in solution to and uniformly distributed on the grains of a suspension agent and dried.
- 25. The method according to any one of claims 19 to 24, wherein the pharmaceutically active substance, before admixing it to the efferwescent system, is mixed with at least one neutral substance, at least one suspension agent and at least one substance selected from the group of alkali carbonate, alkaline auth carbonate, alkaline earth bicarbonate, alkaline auth carbonate, alkaline act bicarbonate, alkaline act bicarbonate, alkaline act of at least one solid edible organic acid, alkaline earth salt of at least one solid edible organic acid, whereafter a solution of at least one binding agent and/or a tenside is at least once applied to, distributed on the grains of the mixture and died.

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- 26. A process for the manufacture of effervescent granules from a powdered or granular mixture of a solid, edible, organic acid and the carbonate and/or bicarbonate of an alkali and/or alkaline earth metal under vacuum, wherein, for the passivation of the sustance of at least one of the components to a state of strong inertia to the reaction, there is added to the heated mixture during the treatment under vacuum a metered quantity of a polar solvent, the difference in pressure caused by development of carbon dioxide through the addition of solvent during the reaction being determined up to a maximum of 1000 mbar, the volume and mass of the carbon dioxide liberated being ascertained from this difference in pressure, and the heat treatment being repeated, after rapid drying of the mixture, as many times as necessary to obtain passivation of the surface as indicated by an evident slowing down of the reaction and by a reduced development of gas, and wherein in said polar solvent there is dissolved a neutral substance selected from the group consisting of a water-soluble polymer, a higher alcohol, a carbohydrate and a hydrocolloid.
- 50 27. A process for the preparation of an effervescent granular material containing at least one solid, crystalline edible organic acid and at least one carbonate of an alkali metal or an alkaline earth metal which splits off CO₂ upon reaction with said organic acid in aqueous solution, which comprises:
- pre-reacting a portion of said organic acid and said carbonate in solution in water and/or alcohol to form a pre-

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adding said pre-reaction product to an additional portion of said organic acid in crystalline form with thorough mixing to form a first coating by reaction with said organic acid crystals and liberation of the resulting water of crystalization,

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- applying at least one additional coating including said carbonate onto the organic acid crystals with said first coating adhering thereto, and
- terminating the reaction after the last coating has been applied by drying, wherein to said pre-reaction product there is added a neutral substance selected from the group consisting of a water-soluble polymer, a higher alcohol, a carbohydrate and a hydrocolloid.

Patentansprüche

- Ein granulares Brauseprodukt, welches zum Herstellen einer wässrigen Lösung oder Suspension einer oder mehrere pharmazeulischa räktver Substanzan zur oralen Verabreichung geeignet ist, und welches in Tableiten pressbar ist, und/oder dieses Produkt in Tableitenform, mit Brausekönrenr, die von Tägerkristallen wentigstens einer lesten, geniessbaren organischen Saure erhalten worden sind, welche Im wesentlichen mit wenigstens einer Beschischtung bedeckt sind, die mindestens eine wassenfosliche, neutrale Substanz enthalt, wobei die neutrale Substanz zum Absenken des Schmelzpunktes der Saurekristalle an ihrer Oberläche wirksam ist, und wenigstens eine eus der aus Alkalicarbonat, Alkalibicarbonat, Erdalkalibicarbonat, einem Alkalisatz und einem Erdalkalisatz zumindest einer festen, geniessbaren organischen Saure bestehenden Gruppe wenigstens einer festen, geniessbaren organischen Staure bestehenden Gruppe wenigstens einer festen, geniessbaren organischen Staure beschichtung angebagd ist.
- Granuláres Produkt oder Tablette nach Anspruch 1, wobei die neutrale Substanz aus der aus einem wasserlösizo chen Polymer, einem höhrenen Alkohol, einem Köhlebydrat und einem Hydrokolloid bestelhenden Gruppe ausgewahlt ist, welche neutrale Substanz in einer Menge von etwa 0,05 bis annahernd 1,0 Gewichts-%, vorzugsweise
 von etwa 0,07 bis ungelähir 0,8 Gewichtle-%, vorhanden ist.
- Granulares Produkt oder Tablette nach Anspruch 1 oder 2, wobei ein Feuchtigkeitsbindentittel an den Brausekörnern verankert ist, welches Feuchtigkeitsbindemittel vorzugsweise aus der aus kalziniertem Soda und Natriumsulfat bestehenden Gruppe ausgewählt ist und vorzugsweise in einer Menge von etwa 4 bis ungelähr 10 Gewichts%, bezogen auf die gesamte Mischung, eingesetzt ist.
- 4. Granulares Produkt oder Tablette nach einem der vonheigehenden Ansprüche, wobei wenigstens eine zusätzliche Beschlichtung an den Brausekörnern angebracht ist, welche eine aus der aus Alkalisalzen und/oder Erdalkalisalzen wenigstens einer festen, geniessbaren organischen Saure bestehenden Grurpe ausgewählte Substanz als Puffer und gegebenenfalls eine zusätzliche neutrale Substanz aufweist, und wobei vorzugsweise wenigstens eine der Beschichtungen ein Antischaummittel enthält.
- Granuläres Produkt oder Tablette nach einem der vorhergehenden Ansprüche, wobei das granuläre Produkt oder das in Tablettenform gepresste granuläre Produkt ferner mindestens ein in einem eigenen granulären Produkt vorhandenes Antischaummittel aufweist.
- Granuläres Produkt oder Täblette nach Anspruch 4 oder 5, wobei das Antischaummittel aus der aus Dimethicon
 und Simethicon bestehenden Gruppe ausgewählt ist und in einer Menge von etwa 0,005 bis ungelähr 0,5
 Gewichts-%, bezogen auf die gesamte Mischung, oder von etwa 0,05 bis ungelähr 2,0 Gewichts-%, bezogen auf
 die pharmazeutisch aktive Substanz, eingesetzt ist.
- Granulares Produkt oder Tablette nach einem der vorhergehenden Ansprüche, wobei es bzw. sie eine Säurebindungsfähligkelt von weniger als 5, vorzugsweise weniger als 3 meq, gemessen nach USP XXII, aufweist.
- Granulares Produkt oder Tablette nach einem der vorhergehenden Ansprüche, wobei es bzw. sie bei einem Gesamtgewicht von nicht mehr als 2,5, vorzugsweise nicht mehr als 2,0 Grannn in Wasser bei Rauntemperatur eine Auflösungszelt von weniger als 180, vorzugsweise weniger als 120, Sekunden aufweist.
- Granulares Produkt oder Tablette nach einem der vorheigehenden Ansprüche, mit einer hydrophoben pliarmazeuitsch aktiven Substanz, wobei die hydrophobe Substanz in von den Bausekomponenten gesonderten Granula vorliegt, in welchen Granula die hydrophobe Substanz auf wenigstens einer aus der aus Suspendiermitten walche vorzugsweise aus der aus Aerosil[®] und Avicel[®] bestehenden Gruppe gewählt sind und neutralen Substanz zen welche vorzugsweise aus der aus Manniol und Sorbiolo bestehenden Gruppe gewählt sind bestehenden Gruppe gewählt sind bestehenden
- 10. Granulares Produkt oder Tablette nach Anspruch 9, wobei die Granula auch wenigstens eine aus der aus Bindern

 vorzugsweise Poływinypyrrolidon (PVP) ., geringen Mengen eines Tensids - welches vorzugsweise aus der aus Diody-Natriumsulfosuozinat und Natriumlauryfsulfat bestehenden Gruppe gewählt ist - , Alkali- und/oder Erdalkaticarbonat und/oder -bicarbonat bestehenden Gruppe ausgewählte Komponente enthalten. 11. Granuláres Produkt oder Tablette nach einem der vorhergehenden Ansprüche, wobei es bzw. sie, bezogen auf die gesante Mischung, etwa 2 bis ungelähr 30 Gewichts-% Cimetidin; wem 30 bis ungelähr 60 Gewichts-% einer lesten, geniessbaren organischen Saure, etwa 12 bis ungelähr 40 Gewichts-% weinigstens eines Aklair-oder Erd alkslicatbonats oder bicarbonats (woron etwa 2 bis ungelähr 10 Gewichts-% kahrumcarbonat als Feuchtigkeits-birdennittel ist); etwa 1 bis ungelähr 4 Gewichts-% eines Süssstolfes; etwa 0,01 bis ungelähr 30 Gewichts-% einer neutralen Substanz (wovon etwa 0,01 bis ungelähr 0,05 Gewichtis-% für die Beschichtung mit neutraler Substanz ist), vorzugsweise etwa 3 bis ungelähr 20 Gewichts-% Sorbitol und etwa 2 bis ungelähr 10 Gewichts-% Mannitol; etwa 0,005 bis ungelähr 0,5 Gewichts-% eines Antischaummittels und etwa 0,1 bis ungelähr 3 Gewichts-% eines Geschinadosnittels.

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12. Granulares Produkt oder Tablette nach einem der Ansprüche 1 bis 10, wobei es bzw. sie, bezogen auf die gesamte Mischung, die folgenden Komponenten enthalt: etwa 0,4 bis ungelähr 4,5 Gewichts-% Binder, vorzugsweise At 4 bis ungelähr 4,5 Gewichts-% Binder, vorzugsweise Polyviny-pyroidon (PVP): etwa 0,03 bis ungelähr 0,35 Gewichts-% Fensid, vorzugsweise Diochyl-Matriumsulfosuccinat; etwa 30 bis ungefähr 56 Gewichts-% wenigstens eines Alkali- undoder Erdalkalicarbonats oder - bicanbonats (wovon etwa 2 bis ungefähr 40 Gewichts-% wenigstens eines Alkali- undoder Erdalkalicarbonats oder - bicanbonats (wovon etwa 2 bis ungefähr 10 Gewichts-% Natriumcarbonat als Feuchtigkeitsbindemittel sind); etwa 0,3 bis ungerens 0,02 bis ungefähr 5,5 Gewichts-% eines Sissstuftes; etwa 0,02 bis ungelähr 0,1 Gewichts-% für die Beschichtung mit neutraler Substanz ist), die vorzugsweise aus der aus Maltodextim, Laktose und Mannitol bestehenden Gruppe ausgewählt ist; etwa 0,005 bis ungefähr 0,05 Gewichts-% eines Antischaummittels, welches vorzugsweise aus der aus Dimethicon und Simetricon bestehenden Gunpa ausgewählt ist; etwa 0,005 bis ungefähr 0,05 Gewichts-% eines Antischaummittels, verless vorzugsweise aus der aus Dimethicon und Simetricon bestehenden Gunpa ausgewählt ist; etwa 0,005 bis ungefähr 0,05 Gewichts-% eines Antischaummittels, verless vorzugsweise aus der aus Dimethicon und Simetricon bestehenden Gunpa ausgewährt ist.

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- Granulåres Produkt oder Tablette nach einem der Ansprüche 1 bis 10, wobei es bzw. sie, bezogen auf die gesamte Mischung, die folgenden Komponenten enthält:
- etwa 0,1 bis ungefähr 0,5 Gewichts-% beta-Carotin (100%);

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- etwa 0 bis ungefähr 2 Gewichts-% Tocopherylazetat (100%);
- etwa 35 bis ungetähr 70 Gewichts % einer festen, geniessbaren organischen Sture, vorzugsweise etwa 0 bis ungetähr 10 Gewichts % Ascotbinsaure, etwa 35 bis ungetähr 55 Gewichts % Zitronensaure und etwa 0 bis ungetähr 5 Gewichts % Maleinsaure;
- etwa 11 bis ungefaltw 38 Gewichts-% wenigstens eines Alkali- oder Erdalkalicarbonats oder -bicarbonats, vorzugsweise etwa 5 bis ungefahr 15 Gewichts-% Calciunicarbonat und etwa 5 bis ungefahr 20 Gewichts-% Natitumbicarbonat;
 - etwa 1 bis ungefähr 4 Gewichts-% eines Süssstoffes;

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- etwa 0,1 bis ungelâhr 35,0 Gewichts % einer neutralen Substanz (wovon etwa 0,1 bis ungelâhr 0,5 Gewichts% für die Beschichtung mit neutraler Substanz ist), die vorzugsweise etwa 1 bis ungelâhr 10 Gewichts-% Sorbitol und etwa 5 bis ungelâhr 25 Gewichts-% Mannitol sind; und
 - etwa 0,3 bis ungefahr 3 Gewichts-% eines Geschmacksmittels
- 14. Granulares Produkt oder Tablette nach einem der Ansprüche 1 bis 10, wobei es bzw. sie, bezogen auf die gesamte Mischung, die folgenden Komponenten enthalt: etwa 3 bis ungelähr 14 Gewichts-% Rantitidin-Hydrochlorid (75 300 mg pro Dosis); etwa 30 bis ungelähr 50 Gewichts-% Natiri-List etwa 2 bis ungelähr 20 Gewichts-% Natiri-List etwa 2 bis ungelähr 20 Gewichts-% Natiri-List etwa 2 bis ungelähr 70 Gewichts-% Natiri-List etwa 2 bis ungelähr 10 Gewichts-% Natiri-List etwa 2 bis ungelähr 0,2 Gewichts-% Natiri-List etwa 2 bis ungelähr 0,2 Gewichts-% einer neutralen Substanz für die erste Beschichtung sowie etwa 0 bis ungelähr 15 Gewichts-% zusatzlicher neutraler Substanzen; etwa 0 bis ungelähr 8 Gewichts-% eines Geschmacksmittels.
- 15. Eine Brauselablette, welche wenigstens eine pharmazeutisch aktive Substanz und ein Brausesystem mit wenigstens siener feiner Istens geniessbaren organischen Säure, wenigstens einem Alkalinetalicarbonat oder -bicarbonat als gasbildende Komponente und mindestens einem Alkalimetalisalz der Säure, wobei zumtndest zwei Schlichten auf Tägerkriställe aufgetirgen sind, welche aus der wenigstens einen Säure bestehen, wobei die erste Schlicht zumindest eine weitere lessle, geniessbare organische Säure oder das Alkalimetalisalz dieser weiteren Säure oder

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beide enthält, wogegen die zweite Schicht mindestens ein Alkalimetallsalz der wenigstens einen Säure enthält, und wobei die erste Schicht zusätzlich eine aus der aus einem wasserlöslichen Polymer, einem höheren Alkohol, einem Köhlehydrat und einem Hydrocolloid bestehenden Gruppe ausgewätilte neutrale Substanz enthält.

- 6 16. Granuläres Produkt oder Tablette mit einem Brausesystem nach einem der Ansprüche 1 bis 15 und Cisaprid als pharmazeubisch aktive Subsianz, wobei, bei einem Gesamtgewicht von weniger als 2 Gramm, vorzugsweise weniger als eiwa 1,6 Gramm, es bzw. sie eine Saurebindungsfähigkeit von weniger als 5 meq, vorzugsweise weniger als 3 meg, besitzt.
- 17. Granuláres Produkt oder Tablette mit einem Brausesystem nach einem der Ansprüche 1 bis 15 und Cimetidin als pharmazeutisch aktive Substanz, wobei, bei einem Gesamtgewicht von weniger als 2,5 Granum, vorzugsweise weniger als etwa 2,0 Granum, es bzw. sie eine Saurebindungsfähigkeit von weniger als 5 meq, vorzugsweise weniger als 3 meq, besitzt.
- 18. Granulares Produkl oder Tablette mit einem Brausesystem nach einem der Ansprüche 1 bis 15 und Ranilldin als pharmazeutisch aktive Substanz, wobei, bei einem Gesamtgewicht von weniger als 2,6 Granum, vorzugsweise weniger als 2,0 Granum, es bzw. sie eine Säurebindungsfaltigkeit von weniger als 3 meq, vorzugsweise weniger als 2 meq, besitzt.
- Verfahren zur Herstellung eines granulären Produktes oder einer Tablette nach einem der vonhergehenden Ansprüche, bei dem Kristalle wenigstens einer festen, geniessbaren organischen Saure mit einer wässrigen i Lösung einer neutralen Substanz angefeuchter wird und dann vor dem vollständigen Trochene ein Alkait- unkfoder Erdaltalfrach bonat undfoder -bicarbonat in Pulverform geichmässig ver leitil und an der feuchten Oberlifachenschicht durch Mischen verankert wird, worauf die so hergestellten Brauselkorner getrodnet und mit einer pharmazeurlisch alkiven Substanz vorzugsweise mit einer staureempfindlichen, insbesondere einer aus der aus H2-Blockern, Cimetidin, Ramitidin, Cisaprid und beta-Carotin bestehenden Gruppe ausgewählten und pharmazeurisch aktzeptablen Hilsmitteln gemischt, und gegebenenfalls zu Tabletten gepresst, werden.
- 20. Verfahren nach Anspruch 19, bei dem auf den Brausekörnern mindestens eine zusätzliche Beschlichtung durch Beleuchten der Körner mit der Lösung einer Puffersubstanz aufgebracht wird, vorzugsweise einer solchen, welche aus der aus Alkelicanbonal, Hanblischsbonal, Erdalkelicanbonal, einem Alkelisalz zumindest einer festen, geniessbaren organischen Saure und einem Erdalkalisalz zumindest einer festen, geniessbaren organischen Saure und einem Erdalkalisalz zumindest einer festen, geniessbaren organischen deutpe ausgewählt ist.
- 21. Verfahren nach Anspruch 19 oder 20, bei dem die Lösung ferner eine aus der aus einem wassenlöslichen Polynier, einem l\u00f6henen Alkohol, einem Kohlehydnat und einem Hydrocolloid bestehenden Gruppe ausgew\u00e4hlt ist.
- 22. Verfahren nach einem der Ansprüche 19 bis 21, bei dem, zusätzlich zum Arzneimittel, die Brausekörner auch mit einem granulären Produkt gemischt werden, das durch Auftragen eines Antischaummittels in einer geeigneten Lösung auf die Oberfläche von Partikeln einer neutralen Substanz hergestellt worden ist, und das Lösungsmittel
- 23. Verfahren nach einem der Ansprüche 19 bis 22, bei dem die getrockneten Brausek\u00f6rner mit \u00e4hnet, das vorzugsweise ein AntlschaummIttel gelöst einhalt, und durch Verdampfen des \u00e4lhanols wieder getrocknet werden, um die Resitauchtigkeit zu beseitigen.

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- 24. Verfahren nach einem der Ansprüche 19 bis 23, bei dem die pharmazeutisch aktive Substanz, vor ihrer Zumischung zum Brausesystem, in Lösung - zusammen mit einem Bindemittel und/oder einem Tensid - auf die Könner eines Suspendlermittels aufgefragen und gleichmässig verteilt und getrocknet wird.
- 25. Verfahren nach einem der Ansprüche 19 bis 24, bei dem die pharmazeulisch aktive Substanz, vor ihrer Zunlischung zum Brausesystem, mit wenigstens einer neutralen Substanz, mindestens einem Buspendiermiltel und zunindest einer aus der aus Alkalicarbonat, Alkalbicarbonat, Erdalkalicarbonat, Erdalkalicarbonat, Erdalkalicarbonat, einem Alkalisat zumindest einer festen, geniessbaren organischen Saure braf besteher Erdalkalisatz zumindest einer festen, geniessbaren organischen Saure besteherden Gruppe ausgewählten Substanz genischt wird, worauf die Lösung werügstens eines Bindemitlets und/oder eines Tensids zumindest einmal auf die Könner der Mischung aufgelagen, verfelt und getrocknet wird.

bestimmt wird, wobei das Volumen und die Masse des freigesetzten Kohlendioxyds aus dieser Druckdifferenz dig ist, um die Passivierung der Oberfläche zu erhalten, wie durch eine deutliche Verlangsamung der Reaktion und geniessbaren organischen Säure und dem Carbonat und/oder Bicarbonat eines Alkali- und/oder Erdalkalimetalls unter Vakuum, bei dem zur Passivierung der Oberfläche wenigstens einer der Komponenten zu einem Zustand starker Trägheit gegenüber der Reaktion der erhitzten Mischung während der Behandlung unter Vakuum eine dosierte Menge eines polaren Lösungsmittels zugefügt wird, die durch die Entwicklung von Kohlendioxyd durch die Zugabe des Lösungsmittels während der Reaktion verursachte Druckdifferenz bis auf ein Maximum von 1000 bar ermittelt wird, und die Wärmebehandlung nach raschem Trocknen der Mischung so oft wiederholt wird, als notwenlöslichen Polymer, einem höheren Alkohol, einem Kohlehydrat und einem Hydrocolloid bestehenden Gruppe aus-Verfahren zur Herstellung von Brausegranula aus einer pulverförmigen oder granulären Mischung einer festen, eine verringerte Gasentwicklung angezeigt wird, und wobei in der polaren Lösung eine aus der aus einem wasser-. 26.

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geniessbare organische Saure und zumindest ein Carbonat eines Alkali- oder Erdalkalimetalls enthält, das bei Reaktion mit der orga-Vertahren zur Herstellung von granufärem Brausematerial, welches mindestens eine feste, nischen Saure in einer wassrigen Lösung CO₂ abgibt, welches folgendes aufweist: 27.

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- vorab Umsetzen eines Teiles der organischen Säure und des Carbonats in einer Lösung in Wasser und/oder Alkohol, um ein Vorreaktionsprodukt zu schaffen,
- Zugeben des Vorreaktionsproduktes zu einem weiteren Teil der organischen Saure in kristalliner Form unter sorgfältigem Mischen, um durch Reaktion mit den Kristallen der organischen Saure und der sich daraus ergebenden Freisetzung von Kristallisationswasser eine erste Beschichtung zu bilden,

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- Aufbringen wenigstens einer weiteren, das Carbonat aufweisenden Beschichtung auf den Kristallen der organischen Säure, an denen die erste Beschichtung anhaftet, und
- Abschliessen der Reaktion, nachdem die letzte Beschichtung aufgetragen worden ist, durch Trocknen, wobei eine aus der aus einem wasserlöslichen Polymer, einem höheren Alkohol, einem Kohlehydrat und einem Hydrocolloid bestehenden Gruppe ausgewählte neutrale Substanz dem Vorreaktionsprodukt hinzugefrigt wird.

Revendications

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d'être pressé en comprimés et/ou ledit produit sous forme de comprimés, comprenant des grains effervescents obtenus à partir de cristaux porteurs d'au moins un acide organique alimentaire et solide, qui sont sensiblement recouverts par au moins un revêtement confenant au moins une substance neutre hydrosoluble, dans lequel ladite substance neutre est capable d'abaisser le point de fusion des oristaux d'acide à leur surtace, et au moins une substance - choisie dans le groupe constitué par les carbonates alcalins, les bicarbonates alcatins, les carbonates alcalino terreux, tes bicarbonates alcalino terreux et les sels alcalins d'au moins un acide organique alimentaire et Produit effervescent granulé, convenant pour la préparation d'une suspension ou d'una solution aqueuse d'una substance active du point de vue pharmaceulique ou davantage, destiné à une administration orale, et susceptible solide et les sels atcalino-terreux d'au moins un acide organique alimentaire et solide - est appliquée sur ledit revêtement. 9

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Produit granulé ou comprimé selon la revendication 1, dans lequel la substance neutre est choisie dans le groupe constitué par les polymères hydrosolubles, les alcools supérieurs, les hydrates de carbone et les hydrocolloïdes et dans lequel ladite substance neutre est présente en une quantité allant d'environ 0,05 à environ 1,0 % en poids et, de préférence, d'environ 0,07 à environ 0,8 % en poids. ٨i

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- Produit granulé ou comprimé selon la revendication 1 ou la revendication 2, dans lequet un agent fixant l'humidité litué par le carbonate de sodium anhydre et le sulfate de sodium anhydre et étant appliqué, de préférence, en une est fixé sur lesdits grains effervescents, cet agent fixant l'humidité étant choisi, de préférence, dans le groupe consquantité allant d'environ 4 à environ 10 % en poids, par rapport au mélange total. က်
- Produit granulé ou comprimé selon l'une quelconque des revendications précédentes, dans lequel on a appliqué sur les grains effervescents au moins un revêtement additionnel, comprenant une substance choisie dans le groupe constitué par les sels alcalins et/ou les sels alcalino-terreux d'au moins un acide organique alimentaire et solide servant de tampon et, à titre facultatif, une substance neutre additionnelle et dans lequel, de préférence, au moins un des revètements contient un agent antimousse. 4
- Produit effervescent granulé ou comprimé selon l'une quelconque des revendications précédentes, dans lequel le 'n

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produit granulé ou ledit produit granulé pressé sous forme de comprimés comprend, en outre, au moins un agent antimousse présent lui-même sous forme d'un produit granulé séparé.

- choisi dans le groupe constitué par la diméthicone et la siméthicone et est appliqué en une quantité d'environ 0,005 à environ 0,5 % en poids par rapport au métange total ou d'environ 0,05 à environ 2,0 % en poids par rapport à la Produit granulé ou comprimé selon la revendication 4 ou la revendication 5, dans lequel l'agent antimousse est substance active du point de vue pharmaceutique. ø,
- Produit granulé ou comprimé selon l'une quelconque des revendications précédentes, ayant une capacité de fixation d'acides inférieure à 5 et, de préférence, inférieure à 3 méq., la détermination étant faite selon USP XXII. ۲. 9
- Produit granulé ou comprimé selon l'une quelconque des revendications précédentes, qui, pour un poids total ne dépassant pas 2,5 et, de préférence 2,0 grammes, présente un temps de dissolution dans l'eau à la température ambiante inférieur à 180 et, de préférence, inférieur à 120 secondes. œ

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- tance active du point de vue pharmaceutique et hydrophobe, et dans lequel la substance hydrophobe est présente dans des granules distincts des composants effervescents, la substance hydrophobe de ces granules étant appliquée en revêtement ou fixée sur au moins une substance choisie dans le groupe constitué par des agents de suspension (choisis, de préférence, dans le groupe constitué par le produit Aerosil ® et le produit Avicel ®) et des Produit granulé ou comprimé selon l'une quelconque des revendications précédentes, qui comprend une subssubstances neutres (choisies, de préférence, dans le groupe constitué par le mannitol et le sorbitol). oi 20
- Produit granulé ou comprimé selon la revendication 9, dans lequel les granules contiennent également au moins un composant choisi dans le groupe constitué par des liants (de préférence la polyvinylpyriolidone (PVPI)), de peti-un composant choisi dans le groupe constitué par des liants (de préférence la polyvinylpyriolidone (PVPI)), de petites quantités d'un tensioactif (choisi, de préférence, dans le groupe constitué par le dioctyl-sullosuccinate de sodium et le fauryf-sulfate de sodium), et les carbonates et/ou les bicarbonates alcalins et/ou alcalino-terreux. 52
- lisé comme agent fixant l'humidité); d'environ 1 à environ 4 % en poids d'un édulcorant; d'environ 0,01 à environ 30 % en poids d'une substance neutre (dont d'environ 0,01 à environ 0,05 % en poids servent pour le revêtement de la substance neutre), de préférence d'environ 3 à environ 20 % en poids de sorbitol et d'environ 2 à environ 10 % 11. Produit granulé ou comprimé selon l'une quelconque des revendications précédentes contenant, par rapport au métange total, d'environ 2 à environ 30 % en poids de cinnétidine; d'environ 30 à environ 60 % en poids d'un acirte organique alimentaire et solide; d'environ 12 à environ 40 % en poids d'au moins un carbonate ou un bicarbonate alcalin ou alcalinoterreux (dont d'environ 2 à environ 10 % en poids sont constitués par le carbonate de sodium ulien poids de mannitol; d'environ 0,005 à environ 0,5 % en poids d'un agent antimousse et d'environ 0,1 à environ 3 % en poids d'un agent aromatisant. 30 35
- total, les composants suivants : d'environ 0,4 à environ 4,5 % en poids de cisapride; d'environ 0,4 à environ 4,5 % en poids d'un agent de suspension; d'environ 0,1 à environ 1 % en poids d'un liant, de préférence la polyvinylpyrrolidone (PVP); d'environ 0,03 à environ 0,35 % en poids d'un tensioactif, de préférence le dioctyl-sulfosuccinate de sodium; d'environ 30 à environ 55 % en poids d'un acide organique alimentaire et solide, de préférence l'acide reux (dont d'environ 2 à environ 10 % en poids sont constitués par le carbonate de sodium utilisé comme agent fixant l'humidité); d'environ 0,3 à environ 2,5 % en poids d'un édulcorant; d'environ 0,02 à environ 55 % en poids d'une substance neutre (dont d'environ 0,02 à environ 0,1 % en poids servent pour le revêtement de la substance neutre), choisie, de préférence, dans le groupe constitué par la maltodextrine, le lactose et le mannitol, d'environ Produit granulé ou comprimé selon l'une quelconque des revendications 1 à 10, contenant, par rapport au mélange citrique; d'environ 12 à environ 40 % en poids d'au moins un carbonate ou un bicarbonate alcalin ou alcalino-ter-0,005 à environ 0,05 % en poids d'agent antimousse, choisi, de préférence, dans le groupe constitué par la dimé thicone et la siméthicone; et d'environ 0,2 à environ 5 % en poids d'un agent aromatisant. 5 9
- Produit granulé ou comprimé selon l'une quelconque des revendications 1 à 10, contenant, par rapport au mélange otal, les composants suivants : 뜫

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- d'environ 0,1 à environ 0,5 % en poids de bêta-carotène (100 %);
- d'environ 0 à environ 2 % en poids d'acétate de tocophérol (100 %);

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d'environ 35 à environ 70 % en poids d'un acide organique alimentaire et solide, de préférence d'environ 0 à environ 10 % en poids d'acide ascorbique, d'environ 35 à environ 55 % en poids d'acide citrique et d'environ 0 à environ 5 % en poids d'acide malique;

- d'environ 11 à environ 38 % en poids d'au moins un carbonate ou un bicarbonate alcalin ou alcalino-terreux, de préférence d'environ 5 à environ 15 % en poids de carbonate de calcium et d'environ 5 à environ 20 % en poids de bicarbonate de sodium;
- d'environ 1 à environ 4 % en poids d'un édulcorant;
- d'environ 0,1 à environ 35,0 % en poids d'une substance neutre (dont d'environ 0,1 à environ 0,5 % en poids servent pour le revêtement de la substance neutre), de préférence d'environ 1 à environ 10 % en poids de sorbitol et d'environ 5 à environ 25 % en poids de mannitol; et
- d'environ 0,3 à environ 3 % en poids d'un agent aromatisant

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- sodique; d'environ 10 à environ 30 % en poids de bicarbonate de sodium; d'environ 2 à environ 10 % en poids de Prothili gravulé ou comprimé selon l'une quelconque des revendications 1 à 10, contenant, par rapport au mélange total, les composants suivants : d'environ 3 à environ 14 % en poids de chlorhydrate de ranitidine (75 - 300 mg par dose); d'environ 30 à environ 50 % en poids d'acide citrique; d'environ 0 à environ 20 % en poids de citrate monocarbonate de sodium; d'environ 1 à environ 3 % en poids d'un édulcorant; d'environ 0,05 à environ 0,2 % en poids de la substance neutre utilisée pour le premier revêtement ainsi que d'environ 0 à environ 15 % en poids de substances neutres additionnelles; d'environ 0 à environ 8 % en poids de granules d'un agent antimousse et d'environ 0,1 à environ 4 % en poids d'un agent aromatisant.
- de métal alcalin en tant que composant générant du gaz et au moins un sel de métal alcalin de l'acide, dans lequel au moins deux couches sont appliquées aux cristaux porteurs constitués par au moins un premier acide, la prevescent comprenant au moins un acide organique alimentaire et solide, au moins un carbonate ou un bicarbonate mière couche contenant au moins un autre acide organique alimentaire et solide ou un sel de métal alcalin de cet autre acide ou les deux, ators que la seconde couche contient au moins un sel de métal alcalin dudit premier acide, la première couche contenant en plus une substance neutre choisie dans le groupe constitué par les polymères Comprimé effervescent contenant au moins une substance active sur le plan pharmaceutique et un système efferhydrosolubles, les alcools supérieurs, les hydrates de carbone et les hydrocolloïdes. ž. 8 33
- cisapride en tant que substance active du point de vue pharmaceulique, qui, pour un poids total de moins de 2 grammes et, de préférence, de moins d'environ 1,6 grammes, a une capacité de fixation d'acides inférieure à 5 16. Protuit granulé ou conpuimé avec un système effervescent selon l'une quelconque des revendications 1 - 15 et du méq. et, de prélérence, inférieure à 3 méq. 30
- Produit granulé ou comprimé avec un système effervescent selon l'une quelconque des revendications 1 15 et de la cimètidine en tant que substance active du point de vue pharmaceutique, qui, pour un poids total de moins de 2,5 grammes et, de préférence, de moins d'environ 2,0 grammes, a une capacité de fixation d'acides inférieure à 5 méq. et, de préférence, inférieure à 3 méq. 7.

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Produit granulé ou comprimé avec un système effervescent selon l'une quelconque des revendications 1 - 15 et de la ranitidine en tant que substance active du point de vue pharmaceulique, qui, pour un poids total de moins de 2,6 grammes et, de préférence, de moins de 2,0 grammes, a une capacité de fixation d'acides inférieure à 3 méq. et, de préférence, inférieure à 2 méq.

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- 19. Procédé de préparation d'un produit granulé ou un comprimé selon l'une quelconque des revendications précédentes, dans lequel des cristaux d'au moins un acide organique atimentaire et solide sont mouillés avec une solution aqueuse d'une substance neutre et ensuite, avant le séchage complet, un carbonate et/ou un bicarbonate alcalin eľou alcalino-terreux, sous forme de poudre, est réparti de manière uniforme et fixé à la couche de surface humide par mélange, suite à quoi les grains effervescents ainsi préparés sont séchés et mélangés avec une substance active du point de vue pharmaceutique - qui est, de préférence, une substance sensible aux acides et qui est choien particulier, dans le groupe constitué par les antagonistes des récepteurs H2, la cimétidine, la ranitidine, le cisapride et le bêta-carotène - et avec des adjuvants acceptables du point de vue pharmaceutique, puis éventuel lement pressés en comprimés. eje.
- Procédé selon la revendication 19, dans lequel on applique sur les grains effervescents au moins un revêtement additionnel, en mouillant les grains avec une solution d'une substance tampon, de préférence choisie dans le groupe constitué par les carbonetes alcalins, les bicarbonates alcalins, les carbonates alcalino-terreux, les bicarbonates atcalino-terreux, les sets alcatins d'au moins un acide organique alimentaire et solide et les sels alcalinoterreux d'au moins un acide organique alimentaire et solide. 20.

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- tance neutre choisie dans le groupe constitué par les polymères hydrosolubles, les alcools supérieurs, les lydrates 21. Procédé selon la revendication 19 ou la revendication 20, dans lequel la solution comprend, en outre, une subsde carbone et les hydrocolloides
- 22. Procédé selon l'une quelconque des revendications 19 à 21, dans lequel, en plus du médicament, les grains eller-vescents sont également mélangés avec un produit granulé qui a été obtenu en appliquant un agent antimousse dans un solvant approprié sur la surface des particules de la substance neutre et en séchant le solvant.
- 23. Procédé selon l'une quelconque des revendications 19 à 22, dans lequel les grains effervescents séchés sont mouillés avec de l'éthanol qui contient, de préférence, un agent antimousse dissout, puis séchés à nouveau, en évaporant l'éthanol, pour enlever l'humidité résiduelle. 9
- 24. Procédé selon l'une quelconque des revendications 19 à 23, dans lequel, ayant de mélanger la substance active du point de vue pharmaceutique au système effervescent, elle est appliquée en solution avec un agent liant evou un tensioactif, et répartie de manière uniforme sur les grains d'un agent de suspension et séchée. 15
- du point de vue pharmaceutique avec le système effervescent, elle est mélangée avec au moins une substance bonates alcalins, les bicarbonates alcalins, les carbonates alcalino-terreux, les bicarbonates alcalino-terreux, les sels alcalins d'au moins un acide organique alimentaire et solide et les sels alcalino-terreux d'au moins un acide Procédé selon l'une quelconque des revendications 19 à 24, dans lequel, avant de mélanger la substance active neutre, au moins un agent de suspension et au moins une substance choisie dans le groupe comprenant les carorganique alimentaire et solide, suite à quoi une solution d'au moins un agent liant eVou d'un tensioactif est appliquée et répartie sur les grains du mélange, qui sont alors séchés. 25 20
- Procédé de fabrication de granules effervescents à partir d'un mélange pulvérulent ou d'un mélange granulé d'un de haute inertie à la réaction, on ajoute au mélange chauffé durant le traitement sous vide, une quantité mesurée d'un solvant polaire, la différence de pression provoquée par la formation de gaz carbonique produit par l'addition du solvant durant la réaction étant choisie pour atteindre au maximum 1000 mbars, le volume et la masse du gaz comme indiqué par un ralentissement évident de la réaction et par une formation diminuée de gaz, une substance neutre choisie dans le groupe constitué par les polymères hydrosolubles, les alcools supérieurs, les hydrates de acide organique alimentaire et solide et d'un carbonate et/ou d'un bicarbonate d'un métal alcalin ou alcalino-terreux sous vide, dans lequel, pour la passivation de la surface d'au moins un des composants pour l'amener dans un état carbonique libéré étant déterminés à partir de cette différence de pression, et on répète le traitement themique après un séchage rapide du mélange, autant de fois que nécessaire pour obtenir une passivation de la surface, carbone et les hydrocolloïdes étant dissoute dans ledit solvant polaire. 26. 52 30 35
- Procédé de préparation d'un matériau granulé effervescent contenant au moins un acide organique alimentaire cristallin et solide et au moins un carbonate d'un métal alcalin ou d'un métal alcalino-terreux produisant du CO₂ par réaction avec ledit acide organique en solution aqueuse, qui comprend les opérations consistant à : 27.
- provoquer une réaction préliminaire d'une portion ducit acide organique et ducit carbonate en solution dans de l'eau et/ou un alcool pour former un produit de réaction préliminaire,

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- talline et procéder à un mélange poussé pour former un premier revêtement par réaction avec lesdits cristaux ajouter ledit produit de réaction préliminaire à une portion additionnelle dudit acide organique sous forne cis-
- appliquer au moins un revêtement additionnel comprenant ledit carbonate sur les cristaux d'acide organique d'acide organique et libération d'eau de cristallisation résultante,

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- terminer la réaction après que le dernier revêtement a été appliqué, par un séchage, une substance neutre choisie dans le groupe constitué par les polymères hydrosolubles, les alcools supérieurs, les hydrates de caravec ledit premier revêtement adhérant à ceux-ci; et
 - bone et les hydrocolloïdes étant ajoutée audit produit de réaction préliminaire.

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(51) Int. CL⁶: A61K 9/46

EUROPEAN PATENT SPECIFICATION

(12)

(45) Date of publication and mention 14.07.1999 Bulletin 1999/28 of the grant of the patent:

(21) Application number: 94203112.1

(22) Date of filing: 26.10.1994

(54) Granular product or tablet containing an effervescent system and an active pharmaceutical substance, as well as a method for its preparation

Ein Brausesystem und einen Arzneiwirkstoff enthaltendes granuläres Produkt bzw. Tablette sowie Verfahren zu deren Herstellung

Procluit granulaire ou comprimé contenant un système effervescent et un agent actif pharmaceutique, et son procédé de préparation

Gergely, Thomas, Dr.

AT BE CHOE DK ES FR GB IE IT LI LUNL PT SE Designated Contracting States: Designated Extension States: <u>8</u>

(30) Priority: 01.03.1994 DE 440664 23.03.1994 CH 87394

Date of publication of application: 06.09.1995 Bulletin 1995/36 43

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Description

vescent system and a - preferably acid-sensitive - pharmaceutical substance, such as cisapride, beta-carotene, an 142 tical preparation with comparatively small amounts of effervescent components or a comparatively low acid-binding (0001) This invention relates to a granular pharmaceutical preparation or more particularly a tablet containing an efferblocker such as cimetidine or ranitidine, and/or a substance which is to be administered in an effervescent pharmaceu-

Background of the Invention

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solution, or in any event, hydrophobic particles of the drug tend to creep upward on the glass. On the other hand, in invention is to provide an effervescent system which will avoid the aforesaid disadvantages and offer the possibility of administering to a patient pharmaceutical substances, inclusive of acid-sensitive substances which have hydrophobic properties or properties influencing the surface tension of water, in pleasant-to-chink effervescent solutions. It is a further object of this invention to create an effervescent tablet or an instant effervescent granular product with an acid bindng capacity of less than 5 meq, in order to avoid undesired antacid effects. This is especially advantageous for all H2 vescent tablets or effervescent instant granular products, since in contact with the acid of the effervescent system such compositions hydrolyze or decompose, i.e. they are not shelf-stable. Furthermore, whenever such a substance also affects the surface tension of water, frothing occurs which is highly undesirable for the consumption of the effervescent certain cases, the antacid side effect of an effervescent tablet is undesirable for many drugs. Therefore an object of this blockers. Lastly, it is desired that the tablet or granular product is to dissolve rapidly in water at a temperature of about [0002] Heretofore it has been possible only with difficulty to incorporate acid-sensitive drugs in stable form into effer 15-20°C in less than about 2 minutes 8

Summary of the Invention

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[0003] The solution to the aforesaid problems can be achieved in a surprisingly simple, cost-effective and efficient manner in accordance with this irvention e.g. by first substantially coating acid particles with a composition comprising at least one neutral substance which causes a depression of the melting point of the acid grains at their surface, and thereafter anchoring thereon at least one second coating which contains an alkali and/or alkaline earth carbonate and/or bicarbonate, and optionally a partial reaction product of the carbonate or bicarbonate with the same or a different 30

[0004] The invention is more fully discussed in detail below along with a detailed discussion and illustration of several preferred embodiments.

Detailed Description

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such as maltodextrin, dextrin and the like; especially preferred are higher alcohols, such as xylitol, mannitol and sorbito under the influence of the only slightly alkaline effervescent-grain surface due to the bicarbonate coating, are subject to Neutral substances within the meaning of this invention include water soluble polymers, such as e.g. polyvinylpyrrolidone, carbohydrates, such as saccharose, pentaerythritol, glucose, and fructose (although the latter two, a Maillard reaction tending to make them yellow and therefore they are not particulary preferred herein), hydrocolloids, Various entoodiments of the invention are described in the defining clauses of the dependent claims. 0005 40

to gluconic acid, can be incorporated at the surface of acid vehicle crystals, with the result that the crystal lattice is distection for acid-sensitive active substances. It has therefore also been impossible hitherto to use the invention of (1006) It is true that W093/00886 discloses that a foreign acid, possibly gluconic acid della-lactone, which hydrolyzes turbed and a melting point depression is achieved. However, such a measure cannot of course provide adequate pro-WO93/00886 for acid-sensitive active substances in practice. 45

a thin polymer layer, such as, for example, with polyvinyl-pyrrolidone, carboxymethylcellulose or the like. However, this results in an undesirable retardation of the dissolution time and, in the case of the 1 to 5% by weight of polyvinylpynoidone described there in the Examples, foam formation problems; furthermore, some acid is always transferred from whereby the acid-sensitive active substances would not be protected sufficiently. In addition, however, those skilled in the art have for over 20 years been unable satisfactorily to solve the problem of accommodating acid-sensitive active ow acid binding capacity and short dissolution time. An effervescent tablet is generally defined as being particularly rapid when the dissolution (or complete suspending) of the tablet components takes less than 120 sec, preferably 90 [0007] It has also been proposed (British Patent 1,270,781) to coat acid vehicle crystals for effervescent tablets with the vehicle crystal to the layer in solution when the coating is applied by means of ethanolic or aqueous solution, substances in effervescent systems not only in a shelf-stable manner but atso in retatively small tablet weights with very 90

notice to the European Patent Office of opposition to the European patent granted. Notice of opposition shall be filed in a written reasoned statement. It shall not be deemed to have been filed until the opposition fee has been paid. (Art. 99(1) European Patent Convention).

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Note: Within nine months from the publication of the mention of the grant of the European patent, any person may give

ac or less

[0008] According to the invention, however, after (preferably only a small amount of) the neutral substance has been applied to the acid grains, alkali and/or earth alkaline carbonate and/or bicarbonate particles are anchored on the grain surface in order to prohibit an interaction between the acid and the active substance.

preserved the process proposed in EP-A1-415 326 for coating acid vehicle crystals with several times the amount of sugar in order, in combination with bicarbonate, to achieve a slightly prickling effect, for a chewable tablet or lozenge has not been able to solve the combination of the problems or tasks: such a system would not be sufficiently reactive to dissolve an effervescent tablet in water within a reasonable time. It was the aim of the said EP-A1 to slow down the reaction between acid and carbonate in order not to produce an undesired high effervescent effect in the mouth.

[0010] It, as disclosed in the prior art (US-A-4 127 645), a tablet having a core of acid, bicarbonate and calcium were coated with a neutral substance, for example with sorbitol solution, such a tablet would not provide reliable protection for acid-sensitive active stotstances contained in the core. However, if the mixture were pressed with a neutral substance (e.g. maltodextrin, if necessary as a mixture with sugar, US-A-4 650 669; sorbitol with vitamins, US-A-5 223 264, only suitable as a pricking chewable table) to give tables, then either both reactants would be coated together or undesisable agglomerated granules would occur. In both cases, the reaction on dissolution of the tablet would take place too slowly and the dissolution time would brus be undesirably increased, or the solution would contain undesirably large amounts of sugar. Furthermore, it is very probable that, in agglomerated granules, acid particles too would be present unprotected at the surface of the granules; however, this results in greater instability for acid-sensitive active substances.

[0011] In U.S. Patent No. 4,867,942, a method is described in which vehicle crystals of a solid, edible organic acid are covered on their surface with a pre-reacted solution serving as buffer, particularly an acid alkali and/or alkaline earth sail of a solid, edible organic acid. Thereafter, more of the acid crystals and amounts of carbonate or bicarbonate are anothored side by side on this coaling. Water which is released in the various neutralization partial reactions is removed by a final treatment with actohol and vacuum drying. Such a process is disadvantageous, however, in that, for acid-sensitive drugs, on the acid crystal surface an additional acid simultaneously enters into a reaction with the alkali carbonate, and the reaction thus proceeds too fast and consequently not sufficiently uniformly. Therefore, the product that forms from this niethod cannot completely prevent the reaction of an acid-sensitive drug mixed in with it, due to the acid crystals tying on the surface of the granules.

(6012) In contrast, the structure of the effervescent system according to this invention not only prevents direct contact of an acid-sensitive droug with the acid crystals thereby providing an effervescent tablet or granular product with substantially more shelf-stability, but it also permits the preparation of substantially smaller tablets, i.e., those with smaller amounts of effervescent components which, when dissolved, result in a buffer system. Thus, the present tablets according to the invention, in contrast to buffer systems of antacid effervescent preparations, can remain at far less than 5 meg of acid binding capacity. Also, in terms of product preparation, a retarded reaction and better compressibility into tablets is obtained. With the aid of this invention, an effervescent tablet can be prepared which for the first time contains an acid-sensitive drup, such as ostapride, or an HZ blocker such as cimetidine, and which has an acid-binding capacity of less than 5 mag for a tablet (or granular product) weight of only 1.6 to 2.3 g.

10013] Further, in accordance with an expectally advantageous embodiment of this invention, after the acid crystals to have been covered with a coordance with an expectally advantageous embodiment of this invention, after the acid crystals intended for a full close can be applied to this coating, so that effervescent grains are formed from acid crystals on which a first coating of neutral substance has formed, and thereon a second coating of carbonate and/or bicarbonate, which has partially reacted with the acid in some cases.

[0014] The invention can be particularly expediently used for products or processes as described, for example, in EP-45 B1-76 340, US-A-4 867 942 and WO93/00886. [0015] The application of the neutral substance, especially a sorbitol solution, for example, causes a depression of the melting point on the surface of the citric acid crystals. Thus, on the one hand, the adhesive force for the next coaling containing point on the surface of the citric acid crystals. Thus, on the one hand, the adhesive force for the next coaling containing ability or alkeline earth carbonates and/or bicarbonates increases, and at the same time this signifies a slower and therefore more uniform reaction of the citric acid crystal surface and better passivation, so that the effervescent grains. On the other hand, the melting point depression profracts the recrystalization time of the citric acid or of the citres that have formed, which signifies better compressibility of the effervescent granules over a longer period of time.

[0016] The amount of neutral substance applied to the acid vehicle crystals depends on the amount of solvent with which the acid can be wetted, since a maximum of 50 - 70 % by weight can be dissolved in an aqueous solution. It is therefore preferably added in an amount of 0.05 bis 1.0, in particular 0.07 bis 0.8, % by weight, based on the acid. Additions of less than 0.07 have only a weak effect and phose of less than 0.07 have no effect which is relevant according to the invention; the shelf-siballity of acid-sensitive acide substances is reduced. Additions of over 0.8 generally begin to have an interfering effect, and at above 1.0 the reactivity of citic acid and of the effervocant system is considerably

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slowed down.

[0017] However, this may be less troublesome in the case of granules since a longer dissolution time tends to be desirable there in order to allow the granules to sink on introduction into water and only thereatter to undergo a reaction for dissolution. Otherwise, however, the amounts of neutral substance which can be applied to, for example, citric acid are determined by the amount of solution with which the citric acid can be wetted, since the neutral substances are in fact applied in solution, and a 50 to max. 70% solution and be prepared. The citric acid crystals cannot be wetted with an infinitely large amount of water and hence solvent.

[0018] In certain droumstances, the neutral coaling, especially if carbonate and/or bicarbonate particles are anchored on it, can also contain small amounts of a solid, edible organic acid, and in some cases an acid different from the one of which the verticle crystals consist; as disclosed per se in another context - but here also in order to intensify the melting point depression and/or to control the effervescent teaction and rate of dissolution.

[0019] Each such effervescent grain is, taken by itself, actually a small effervescent "tablet", and effervesces by itself

[0020] Therefore, if desired, a short dissolving time, small quantity and low acid-binding capacity can be achieved. [0021] Experiments thus far towards achieving a fast-acting, small effervescent tablet by the use of monosodium oils rate instead of ctiric acid have failed, because this greatly slows the effervescent reaction, since the monosodium citate reacts more slowly with sodium bicarbonate, and such tablets usually have an acid consuming capacity exceeding 5

[0022] On the other hand, a very thin monosodium citrate coating in accordance with this invention, especially as a third of fourth layer, which can contain an additional neutrial substance if desired, acts advantageously because 1 mol of monosodium citrate binds 1 mol of water of crystallization and thus contributes to the drying or to maintenance of dryness. Furthermore, in any case, uncovered cittic acid surfaces can be covered again or more completely with bicarbonate.

[0023] Additionally, since many substances exhibit some form of taste sensation of which many are unpleasant, especially those exhibiting bitterness, it is desirable to keep the final effervescent solution, especially since it is in beverage 5 form, within the pH range of 3.8 to 4.6. Experience has shown that within this range paricularly bitter substances can be more effectively masked.

be more effectively masked.

10024] While not obligatory, it is preferable to remove residual water from the reaction granules in the course of their preparation by a final treatment with alcohol. Alcohol may disrupt the bonding of water of drystalization, because during drying the residual moisture is removed along with the alcohol by evaporation. Small amounts of an antitioaming agent as on also be added to the alcohol in order to accelerate the dissolution of the final tablet.

[0025] Many of the alorementioned ducy, especially cinetidine and disaptide, other cause frollning in an effervescent tablet. This is not due, however, to foaning such as that caused by tensides. That is to say, the active agents then selves, when stirred into water, do not foam. Instead, when the effervescent particles in the tablet dissolve, butbles of carbon dioxide form.

19026] These bubbles burst and leave the CO₂ on the surface. Now, if a less soluble or more hydrophobic substance is present, the undissolved particles envelop the CO₂ bubbles, and by forming such at film successfully prevent rapid bubble bursing, so that the bubbles with this film on the surface collect and thus a "leam" is formed. However, the "loam" already forming between the effervescent grains interferes with the continued reaction, and thus with the rapid dissolution of the tablet or granules. This circumstance is combatted according to the invention by the addition of very small amounts of at least one antifoaming agent with the result that any "loam" that forms as the effervescent reaction

begins immediately collapses.

[0027] The antibanning agent is preferably added in an amount of 0.005 to 0.5% by weight, based on the total amount including any fillers, flavors, etc., or 0.05 - 2.0% by weight, based on active substance. Additions of less than 0.005 have no effect relevant according to the invention; additions of more than 0.5 give rise to troublesome or unacceptable side effects.

[0028] In the case of active substances which are soluble, although not too freely soluble, as in the case of chrelifdine, a percentage of simelihizone of 0.1 o.03% by weight, based on active substance, is used, which is equivalent to the use of 0.016 - 0.028 percent (about 0.03%) based on the total tablet weight. The situation is somewhat different in the case of an insoluble hydophobic active substance, such as cisapride (the monohydrate is used), where 1% of simelihizone is used, based on the active substance, but an amount of 0.006% results when based on the tablet weight of 1.6 g. It is evident that the disapractie, as a slightly soluble, hydrophobic active substance, requires a larger amount of antificaming agent for suppressing the foam, but the required fillers and the effervescent base result in a substantially smaller amount of simethicone being used per tablet, so that the ritery are inverted.

[0029] In the case of the soluble active substances, such as cimetidine and ranitidine, the simethicone is required in smaller amounts, in order to suppress the smaller landency to loaming in the local reaction on dissolution of the effer-vescent tablet, whereas in the case of cisapiride - as already mentioned - the tendency to foam is substantially greater and the principle is therefore also slightly different.

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0030) If larger amounts are used, film formation of simethicone occurs at the surface after dissolution of the efferves-

stance collect and remain hanging and thus result in unattractive dissolution behavior, this film then additionally having cent tablet, by virtue of the fact that - especially in the case of insoluble active substances - particles of the active subthe tendency to form a ring on the glass wall.

- (9031) In some cases, however, very small amounts of a tenside, for example, docusate sodium, are also added. Due to their wettable nature, such drug particles dissolve more quickly and no longer adhere to the foam bubbles. The proportion of such substances must be determined very precisely to achieve the desired dissolving characteristics.
- ment of the desired effect are used. In the second case, only those drugs are involved which, when the antifoaming of their solubility or stability. Additionally, in the course of production with the use of finely powdered drugs the addition of antitioaming agents may lead to poor distribution because of drug particles attaching themselves to the antifoaming (0032) Although in some cases the antifoaming agent can be applied to the effervescent system and/or to the drug, agent is drawn onto them from a solution in a solvent (e.g., methyl ethyl ketone and acetone) at 40°C, do not lose any this is not preferred according to the invention. In the first case, it might cause undesirable slowing of the dissolution and reaction of the effervescent components unless very slight amounts of antifoaming agent sufficient for the achieveagent droplets.
- It is therefore preferred, in accordance with this invention, that first the formation of a typical granular product irom antifoaming agents and a neutral substance is undertaken, which product is thereafter mixed with the effervescent system and the drug, plus additional adjuvants if desired (e.g., perfumes, sweeteners and the like) and the mixture then compressed into tablet form.
- best be bound by the addition of a moisture-binding agent, especially anhydrous sodium carbonate (which can absorb The moisture released in the preparation of the effervescent system by the neutralization reaction, and not entirely removed by heating and/or vacuum treatment, as well as moisture picked up from the air during storage, can 10 mols of water per mol) or sodium sulfate. The agent can be bound either by applying it to one or more of the coatings applied to the vehicle crystals, or by adding it to the total mixture. This improves shelf life because the reaction of the However, excessive amounts of such moisture-binding agent, for example sodium carbonate, are not desirable as it acid-sensitive active agent with the acid is further suppressed or completely prevented by the reduction of moisture. [0034]
 - Sodium carbonate as a drying agent, therefore, should not be used for completely covering the effervescent may retard the effervescent reaction. [0035]
- grains, since it is preferable to operate with only small quantities effective to merely dry the residual moisture, or to relard the reaction during manufacture, and to avoid undesirably lengthening the dissolving time of the tablet. Therefore, the final addition of sodium carbonate should not be used for complete coverage (or a tablet coating), due to both the quantity and the grain size (approx. 0.1 - 0.05 mm), and it is therefore not suitable for producing a continuous coating on the bicarbonate already present. However, it can be partially hooked onto the effervescent grains. It is also possible, however, not to add the sodium carbonate until after the drying operation. 30
 - In principle, the percentage amount of sodium carbonate per tablet depends on several factors, such as, for exantole, the antount of effervescent base used, the amount and type of the fillers used, the presence of other carbonates, such as, for example, calcium carbonate, etc. [0036] 35
- and 10, in particular 4 6, % by weight (based on the total amount, including any fillers, flavors, etc.). Additions of tess have no effect relevant according to the invention. Additions of over 6 generally begin to have a troublesome effect because sodium carbonate dissolves more slowly and reacts more poorly; above 10% the dissolution time is already [0037] The moisture-binding agent, in particular sodium carbonate, is preferably added in an amount of between 1 than 4 have only a weak effect, and with those of less than 1, the drying effect and increase in stability is too small, they significantly lengthened, since sodium carbonate first absorbs water (up to 10 molecules of water of crystallization) on dissolution of the effervescent tablet, i.e. is calcined and only then reacted with the citric acid. 9
 - 10038] Here it is to be emphasized that 1 mol of water of crystallization can be bound per mol by sodium citrate alone developing in or on the scribitol layer, and in spite of any residual moisture present the scribitol layer prevents or hinders any acid harm to the drug. 5
- If all of the prescribed steps are followed in accordance with the invention, effervescent tablets can be produced, even with the difficult substances referred to, which at a tablet weight of, e.g., 1.6 g.will attain a dissolving time of less than 100 seconds. It is also to be noted that especially cimetidine, due to its hydrophobic character, further lengthens the dissolving time in comparison with other drugs, under otherwise equal conditions. [0039] ŝ
- [0040] Granulation with sorbitol solution permits rapid dissolution without the incorporation of an extraneous acid that is otherwise necessary, for example, according to WO93/00886.
 - Furthermore, during the preparation of the effervescent systems of this invention, and in any case of the tablets themselves, the steps taken according to the invention will enable the control of reactions which take place at the surlace of individual crystals or granules, which thus constitute a local mechanism, while also during dissolution the abovedescribed desired advantages will be achieved throughout. [0041] 55
- (0042) The system is also extraordinarily well suited for the processing of substances which are both acid-sensitive and sparingly soluble in water. Such substances, such as cisapride for example, exhibit very unpleasant behavior in

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suspension, since, as mentioned above, they tend to froth together with the effervescent system, adhere to a glass wall, form unpleasant rings and tend to aggiomerate on the surface of the drink.

- 0043 All the aforesaid problems can be effectively combatted by preparing seperate granules. For this purpose in yet another embodiment of this invention, there is provided a vehicle which can consist of an Aerosi and/or a neutral substance, on which the drug is applied preferably with the surface of its grains partially dissolved, and/or will binding agents and/or tensides it desired, and dried, or is bound to the vehicle surface by means of binders.
- 10044] The amount of the suspended substance is limited to at most 8, preferably at most 4.5, % by weight (based on since it otherwise leads to undesirable agglomerated granules of active substance, suspended substance and binder, the total mixture), for example for cisapride, since larger amounts would result in increased sinking of the granule par ticles after dissolution of the tablet. On the other hand, the amount of binder used is likewise limited to 1% by weight which dissolve only with difficulty and then sink to the bottom, i.e. prevent the desired suspension. 2
- [0045] The invention will now be more fully described and understood with reference to the following examples of preferred embodiments.
 - [0046] Alternatively, the drug can also be dissolved in the methyl ellyl ketone or in acetone and coated onto mannitol Aerosil(R) and sodium bicarbonate. 5

Example 1: Preparation of effervescent tablets containing 200 mg of cimetidine

a) Preparation of the effervescent system

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face on which up to about 30% of bicarbonate can be anchored) or tartaric acid are aspirated into a preheated vacuum tank and heated to approx. 60°C with stirring. Next, 0.85 parts by weight of a solution 1, which has been formed from nate, is aspirated and distributed on the citric acid by mixing. Thereafter, 52.5 parts by weight of sodium bicarbonate and 4.4 parts by weight of aspartame are added to this mixture, which is then stirred and dried by a vacuum of up to erable for improving build-up to effervescent grains on the vehicle crystal as the powder particles provide a rough sur-36 parts by weight each of water and sorbitol, 21 parts by weight of citric acid and 7 parts by weight of sodium bicarbo-200 mbar, after which 1.9 parts by weight of sodium carbonate are aspirated and distributed in the mixture by stirring. 102 parts by weight of coarse citric acid and 25 parts by weight of finely powdered citric acid (the latter is prefand the mixture is then dried by a vacuum of up to 15 mbar.

10048] Next, a further 0.6 parts by weight of said solution are aspirated and distributed in the mixture by mixing. The resultant effervescent grains are dried in a vacuum of up to 20 mbar with stiming. If necessary, 0.25 parts by weight of 96% ethanol are also employed to dry the mixture, and aspirated. Then, again 9.3 parts by weight of sodium carbonate are bound onto the effervescent grain surface. After another final drying, the product is removed through a sieve. 30

b) Preparation of the granulated antifoaming agent 35

In a vacuum mixing tank with a jacket temperature of 80°C, 7.7 parts by weight of sorbitol powder are added and heated to 50°C Then, 0.2 parts by weight of simethicone in a 30% butanone/acetone mixture (5:3) are aspirated in, slirred by vibrational mixing and dried under full vacuum down to 15 mbar at a temperature of at least 45°C.

c) Preparation of the total mixture

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for 10 minutes at 6 rpm with 178.4 parts by weight of the effervescent system prepared in a). Then 7 parts by weight of the antitoanting agent granules prepared in b) and screened through a 0.6 mm sieve, and 4.5 parts by weight of lemon flavoring, are added, mixed for another 5 minutes at 6 rpm. The final mixture is pressed into tablets which weigh 2.3 g. In a mixer, 20 parts by weight of cimetidine, with 21.1 parts by weight of sorbitol powder if desired, are mixed contain 200 mg of cimetidine, and have a hardness of 6-8 kp. [0000] 45

Example 2: Preparation of effervescent tablets containing 200 mg of cimetidine, and citric and malic acid in the effervescent grains: 20

102 parts by weight of coarse citric acid, 25 parts by weight of powdered citric acid and 1.1 parts by weight of malic acid are heated to 60°C with stirring in a preheated vacuum tank. A solution consisting of 0.4 parts by weight of water, 0.22 parts by weight of sorbitol and 0.22 parts by weight of malic acid is then aspirated in and distributed onto the citric acid by mixing. 52.5 parts by weight of sodium bicarbonate and 4.4 parts by weight of aspartame are next added to the mixture and dried by stirring, in a vacuum of up to 200 mbar. Next, 1.9 parts by weight of sodium carbonate are aspirated in and distributed in the mixture by stirring, and then vacuum drying is performed down to 15 mbar. Finally, a final drying can be performed with ethanol, and then 9.3 parts by weight of sodium carbonate are added to the mix-55

ture. The rest of the procedure is similar to Example 1.

Example 3: Effervescent tablets containing 400 mg of cimetidine, and mannitol as a neutral substance

[0052] 49 parts by weight of citic acid are aspirated into a preheated vacuum tank and heated with stinring to 60°C. Then, 0.45 parts by weight of a solution 1, which has been prepared from 0.25 parts by weight of water and 0.20 parts by weight of a solution 1, which has been prepared from 0.25 parts by weight of susparated in and distributed on the citic acid by mixing, whereupon 14.7 parts by weight of social more acid by mixing, whereupon 14.7 parts by weight of social more by weight of asparated and distributed on the citic acid by mixing, whereupon 14.7 parts by weight of social more acid more acid and distributed and distributed by mixing is performed with a vacuum to 15 mbar. Then 0.5 parts by weight of a solution 2, which has been prepared from solution 1 by the addition of 0.16 parts by weight of monosodium citrate, is aspirated into the mixture and distributed by mixing. The effervescent grains obtained therefrom are then died by vacuum and stinring to 2 mbar, and finally 2.8 parts by weight of social more are added. To this mixture are then added 17.3 parts by weight of antitioaming agent granules prepared according to Example 1 b), until distribution is uniform.

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Example 4: Effervescent tablets containing 300 mg of cimetidine, as well as maltodextrin as a neutral substance

20 [0053] Similarly to Example 3, for a 300 mg cimetidine effervescent tablet, a 50% solution of maltodextrin is selected, which is then used in the same amount as in the case of the 400 milligram form.

[0054] In all of the examples in which the tablets contain 100 to 400 mg of cimetidine, the tablet weight can be 2.3 g. The tablets have a dissolving time of preferably 60 to 150 seconds and a buffering capacity below 5 meq, measured according to USP XXII, by back-titration (with 0.5 N NaOH) of an effervescent tablet dissolved in 70 ml of water and with 30 ml of 1.0 N HCl added.

[0055] The figures given in the following table 1 are the percentages of individual ingredients in the particular total mixture of the illustrated preferred embodiments, which therefore are in the following preferred ranges:

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Table 1

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Cimetidine	2 - 30%	(corresponds tablet containir cim	(corresponds to an effervescent tablet containing 50 to 600 mg of cimetidine)
Citric acid	30 - 60%	sorbital	5-20%
Sodium bicarbonate	10 - 30%	mannitol	2-10%
Sodium carbonate	2 - 10%	simethicone	0.005-0.5%
Aspartame	1-4%	flavoring	0.1-3%

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[0056] A preferred percentage composition of cimetidine effervescent tablets or bags of granules containing 100, 200, 300 and 400 mg of cimetidine, with a total weight of 2.31 grans, is summarized below in table 2.

Table 2

	100 mg	200 mg	300 mg	400 mg
Cimetidine	4.30	8.70	13	17.3
Citric acid	20	20	48.2	48.2
Sodium citrate	0.04	0.04	0.04	0.04
Aspartame	1.74	1.64	2.54	3.24
Sorbitol	12.5	12.5	12.8	8.00
Sodium bicarbonate	20.7	20.7	14.7	14.7
Sodium carbonate	4.4	4.4	3.5	3.3
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Table 2 (continued)

	100 mg	200 mg	300 mg	400 mg
Manntiol	4.3		4.3	4.3
HMA Lemon flavoring	2.0	2.0	6.0	6.0
Simethicone	0.05	0.02	0.05	0.05

Example 5: Cisapride effervescent tablets

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a) Preparation of the effervescent grains

[0057] 655 parts by weight of crystalline citric acid, 70 parts by weight of citric acid powder and 8 parts by weight of sodium are heated while mixing to 60°C. Then 2.8 parts by weight of a solution consisting of 0.6 parts by weight of a solution consisting of 0.6 parts by weight of sorbids, 0.3 parts by weight of this cacid and 1.6 parts by weight of water are aspirated into this mixture and distributed by mixing. Next, 340 parts by weight of sodium bicatbonate as well as 2 parts by weight of aspartame are added and reached. Before drying, 77 parts by weight of sodium carbonate are added, whereupon the mixture is vacuum dried with slow stirring to 15 mbar.

20 b) Preparation of the granulated drug

[0058] Insoluble and hydrophobic disapride is attached to a suspending substance by means of a binder and a small amount of a tenside as follows: A solution of 10 parts by weight of disapride, 2 parts by weight of polyvinylpyriolidone and 0.8 part by weight of docusate sodium in 1 part by weight of ethanol and 40 parts by weight of acetone is applied to 10 parts by weight of Aerosil⁽⁴⁾, uniformly distributed and then dried white stirring. The granules are sieved to 0.1.

c) Preparation of the end mixture

30 [0059] To 1152 parts by weight of effervescent grains are added 50 parts by weight of maltodextin, 100 parts by weight of lactose, 184 parts by weight of mannitol, 40 parts by weight of flavoring, 50.2 parts by weight of anti-toanning granules (0.2 parts by weight of simethicone coaled onto 50 parts by weight of mannitol), as well as the granulated drug prepared in b), nixing is carried out for 15 minutes for uniform distribution and the mixture is then pressed to form tablets of 1.6 g, which have an acid-dading capacity of only 2 meq. Gisapride effervescent tablets having such a low acid- 35 binding capacity are unknown to date.

Example 6: Beta-carotene effervescent tablets

[0060] With this extremely acid- and oxidation-sensitive aubstance, attention must be paid to an especially good covering of the acid. The surface and the contact zone on the beta-carotene must be kept alkaline. Therefore the efferves cent grains are covered at least in part with calcium carbonale, thus insuring an alkaline surface. This, however, does result in a slightly longer dissolving time, which in this case is desirable, because the beta-carotene needs time to suspend while the tablet is dissolving. Large amounts of sorbitol, as in US-A5 223 264 mentioned at the outset, are by no means suitable for a beta-carotene effervescent tablet which is intended to be dissolved or suspended in water.

a) Preparation of the effervescent grains

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[0061] 1315 parts by weight of citric acid. 7 parts by weight of sodium saccharin and 45 parts by weight of sodium cyclamate are heated in a vacuum lank to 50°C. Then 16.8 parts by weight of a solution prepared from 3.6 parts by everyth of calcium carbonate, 19 parts by weight of carbon land is a solution and 5 parts by weight of weight of weight of carbon and of shibuted onto the citric acid by mixing. Next, 400 parts by weight of selection carbonate and 190 parts by weight of ordic acid are added and the mixture heated with stirring to 60°C. Then follows the second granulation with 44 parts by weight of the above-mentioned solution, and after distributing and mixing, 403 parts by weight of socium bicarbonate are added, and also, before drying, 52 parts by weight of socium carbonate. The mixture is then second granulation with 45 parts by weight of solver drying, 52 parts by weight of socium carbonate. The mixture is then

b) Preparation of the end mixture

130 parts by weight of sorbitol and 540 parts by weight of mannitol and 50 parts by weight of flavoring, an encapsulated beta-carotene suspendable in water and corresponding to 2 to 15 parts by weight of 100% beta-carotene, plus, it desired, 50 to 250 parts by weight of vitamin C and/or a solid tocopheryl acetate suspendable in water (corresponding to 10 to 75 parts by weight of 100% tocopheryl acetate), plus still other vitamins if desired, are mixed with 2415 parts by weight of the effervescent grains prepared according to a). The product has a tablet weight of 3.3 g and its dissolving time is 60 to 90 seconds.

Example 7: Ranitidine effervescent tablets 5

a) Preparation of the effervescent grains

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840 parts by weight of crystalline citric acid, 210 parts by weight of citric acid powder, 45 parts by weight of sodium cyclamate, and 4 parts by weight of sodium saccharin are heated in a vacuum mixing tank at 60°C. Then a sotuion consisiing of 6 parts by weight of water, 1 part by weight of sodium citrate, and 3 parts by weight of sorbitol is aspithereafter 370 parts by weight of monosodium dirate are added, which are also allowed to react. Lastly, 100 parts by rated in and distributed by stirring. 500 parts by weight of sodium bicarbonate are next added and allowed to react, and weight of sodium carbonate are added and the granules are dried with slow stirring up to 15 mbar. [0063]

b) Preparation of the end mixture

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of mannitol plus 100.4 parts by weight of a granulated antifoaming agent (consisting of 100 parts by weight of mannitol and 0.4 parts by weight of simethicone) and the flavoring agent are added. This mixture is mixed for 15 minutes for uniform distribution, and then pressed to tablets of 2.5 g. The tablets have a dissolving time of 60 to 80 seconds and an To the effervescent grains thus prepared, 167 parts by weight of ranitidine hydrochloride, 125 parts by weight acid-binding capacity of about 2 meq and contain (in percent by weight) 6.8 ramitidine hydrochloride, 42.0 citric acid, 14.8 monoscotium citrate, 20.0 sodium bicarbonate, 4.0 sodium carbonate, 2.0 sweeteners, 5.0 mannitol, 0.1 sorbitol, 4.0 granulated antifoaming agent (containing 0.016 diemthylpolysiloxane) and 1.2 flavoring.

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Example 8:

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545 parts by weight of crystalline citric acid and 133 parts by weight of powdered citric or tartaric acid are parts by weight of sorbitol is distributed on the surface by stirring. Next, 222 parts by weight of sodium bicarbonate are parts by weight of ranitidine hydrochloride, 100 parts by weight of anti-foaming granules (containing 0.4 parts by weight of simethicone and 100 parts by weight of lactose), plus 54 parts by weight of sweetener and 40 parts by weight of flavoring, until unitorm distribution is obtained. The mixture is then pressed to tablets weighing 1.43 g and having a dissolving time of 65-70 sec, a hardness of 8 kp, and an acid-binding capacity of about 1.5 meq. The product contains no monosodium citrate. Ranitidine effervescent tablets having such a low acid-binding capacity have not been disclosed rnixed white heating to 60°C. Then, as the first coating, a solution which consists of 6 parts by weight of water and 4 made to react on the surface of the citric acid, and finally 80 parts by weight of sodium bicarbonate are added. The product is dried with slow stirring. The granules are screened to 1.5 mm, and then mixed for 10 minutes at 10 rpm with 167 [900] to date. ç

Example 9: 45

[0066] 38 2% of citic acid is heated with 0.26% of sodium saccharin to 60°C, then two-thirds of a solution which consists of, with respect to the final mixture, 0.6% water, 0.18% sorbitol, and 0.12% sodium citrate are applied. The solution is distributed for 5 minutes by mixing at 10 rpm. Then 16.2% of sodium bicarbonate and 2.9% of aspartame are added and anchored on the surface of the citric acid by reaction on the neutral substance coating. Then follows a second wetting with the third one-third of the solution; then 12.9% monosodium citrate and, tinally, 5.2% sodium carbonate are added. The effervescent grains are dried while mixing them slowly by applying a vacuum, at a temperature of at least 50°C, to 15 mbar. The basic effervescent granular product is screened to 1.5 mm and mixed with 11.0% of ranitidine hydrochloride, 6.5% of mannitol, 6.5% of anti-foaming granules plus 0.2% of flavoring, and pressed to tablets of 1.55 g, which have a dissolving time of 50 sec at a hardness of 7.3 kp and an acid-binding capacity of less than 2 meq. 20 ß

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Example 10: Vehicle crystal grains coated only with a neutral substance

Since cisapride, for example, in comparison to ranitidine, is not as highly sensitive to acid, it is nevertheless also possible by the procedure to be described below to achieve protection against the acid, all the more so since the drug is embedded in granules. [0067]

a) Preparation of the acid crystals coated with a neutral substance

Then a solution of 4 parts by weight of sorbitol in 4 parts by weight of water is applied and distributed onto the surface 593 parts by weight of crystalline citric acid plus 70 parts by weight of citric acid powder are heated to 60°C of the citric acid by mixing. Finally the citric acid thus coated is vacuum dried at 50 to 60°C. [0068] 2

[0069] In the case of both the form of effervescent product presented here and that of effervescent grains which contain a second alkali or alkali earth carbonate coating, it is possible to protect cisapride, for example, against attack by the citric acid in the drug granules by the addition of sodium bicarbonate.

b) Preparation of the drug granules

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[0070] 160 parts by weight of mannitol, 10 parts by weight of cisapride, 5 parts by weight of aerosil and 10 parts by weight of sodium bicarbonate are heated with mixing to 60°C. Then half of a solution of 27 parts by weight of methyl ethyl ketone (or 45 parts by weight of acetone), 2 parts by weight of alcohol, 2 parts by weight of poly(vinyl pyrrolidone) K30, 1 part by weight of propylene glycol and 0.8 parts by weight of docusate sodium are added and distributed for 5 minutes for the purpose of uniform wetting. The mixture is dried to 0.8 bar, the second part of this solution is aspirated, and again distributed by stirring for 5-10 minutes, and finally vacuum dried. 20

[0071] The active agent granules are then screened to 0.3 mm and already have an enhanced protection against acid attack simply due to the sodium bicarbonate they contain. They can then be mixed with the acid crystals coated with neutral substance, the remaining carbonates and bicarbonates, as well as the other tablet ingredients, and pressed to give tablets. 52

c) Preparation of the end mixture

The citric acid dried and coated according to a) is then mixed with the drug granules prepared according to t), 50 parts by weight of sweetener, 80 parts by weight of sodium carbonate, 430 parts by weight of sodium bicarbonate, and 50 parts by weight of maltodextrin, 100 parts by weight of lactose, 150 parts by weight of mannitol, 50 parts by weight of an antifoaming granulate, and 20 parts by weight of flavoring, and then pressed to tablets of about 1.6 g, which have a dissolving time of 60 to 70 seconds at a hardness of 7 kp. 30 35

Example 11: Cisapride effervescent tablets

a) Preparation of the effervescent granules:

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40 parts by weight of powder, together with 5 parts by weight of saccharin sodium, is uniformly wet at 60°C with 2.2 0.45 part by weight of citric acid and 1.2 parts by weight of water. 12 parts by weight of malic acid are then aspirated in and uniformly anchored on the sorbitol layer formed on the citric acid crystals. Finally, 205 parts by weight of sodium bicarbonate and 1.2 parts by weight of aspartame are aspirated in and once again uniformly distributed. Finally, the Citric acid, consisting of an amount of 300 parts by weight of granules, 80 parts by weight of fine granules and parts by weight of a solution which contains 0.4 part by weight of sorbitol, 0.15 part by weight of sodium bicarbonate, material is covered with 46 parts by weight of sodium carbonate, vacuum dried and discharged through a 1.2 nim sieve. 100731 5

b) Preparation of the active ingredient granules:

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[0074] 12 parts by weight of polyvinylpyrrolidone are dissolved in 12 parts by weight of ethanol; 6 parts by weight of propylene glycol and 6 parts by weight of docusate sodium are then added and the mixture is diluted with 165 parts by weight of ethyl methyl ketone. Half of this solution is distributed over a mixture of 960 parts by weight of mannitol, 30 parts by weight of Aerosii^(f), 60 parts by weight of sodium bicarbonate and 61 parts by weight of cisapride, which is heated to 60°C. Partial drying is then carried out in vacuo, and further wetting is effected with the second half of the solution, followed by complete drying and dicharge through a 0.3 mm sieve. 55

The end mixture is prepared analogously to Example 5.

Claims

- 1. A granular effervescent product suitable for preparing an aqueous solution or suspension of one or more pharmaceutically active substances for oral administration, being equable to being pressed into tablets, and/or said product in label form, comprising effervescent grains obtained from earrier crystals of at least one solid, edible organic acid which are substantially covered by at least one coating containing at least one water-soluble neutral substance, wherein said neutral substance is effective for depressing the melting point of the acid crystals on their surface, and at least one substance selected from the group consisting of alkali carbonate, alkali bicarbonate, alkaline earth hisarbonate, alkaline earth bicarbonate, alkali sail of at least one solid edible organic acid; alkaline earth sail of at least one solid edible organic acid; alkaline earth sail of at least one solid edible organic acid; alkaline earth sail of at least
- The granular product or tablet according to claim 1, wherein the neutral substance is selected from the group consisting of a water-soluble polymer, a higher alcohol, a carbohydrate and a hydrocolloid, and which neutral substance is present in an amount of from about 0.05 to about 1.0 % by weight, preferably from about 0.07 to about 0.8 % by weight.

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- The granular product or tablet according to daim 1 or 2, wherein a moisture-binding agent is anchored on said
 effervescent grains, which moisture-binding agent preferably is selected from the group consisting of anhydrous
 sodium carbonate and sodium sulfate and preferably is applied in an amount of from about 4 to about 10 % by
 weight with respect to the total mixture.
- 4. The granular product or tablet according to any one of the preceding claims, wherein on the effervescent grains at least one additional coating is applied, comprising a substance selected from the group consisting of alkelit salts and/or alkaline earth salts of a teast one solid, edible, organic acid as buffer and, optionally, comprising an additional neutral substance, and wherein preferably at least one of the coatings contains an antillorning agent.
- The granular effervescent product or tablet according to any one of the preceding claims, wherein the granular product, or said granular product compressed in tablet form, further comprises at least one antifoaming agent present in a granular product of its own.
- The granular product or tablet according to daim 4 or 5, wherein the antitioaming agent is selected from the group
 consisting of dimethicone and simethicone and is applied in an amount of from about 0.005 to about 0.5 % by
 weight with respect to the total mixture or from about 0.05 to about 2.0 % by weight with respect to the pharmaceutically active substance.
- The granular product or tablet according to any one of the preceding claims, wherein it has an acid-binding capacity of less than 5, preferably less than 3 meq, measured according to USP XXII.
- The granular product or tablet according to any one of the preceding claims, wherein, at a total weight of no more
 than 2.5, preferably no more than 2.0 grams, in water at room temperature, it has a dissolving time of less than
 180, preferably less than 120 seconds.
- 9. The granular product or tablet according to any one of the preceding daims, comprising a pharmaceutically active substance which is hydrophobic and wherein the hydrophoblc substance is present in granules separate from the effer vescent components, in which granules the hydrophobic substance is coated or anchored onto at least one substance selected from the group consisting of suspending agents preferably selected from the group consisting of derosifing agents preferably selected from the group consisting of derosifing agents preferably selected from the group consisting of accounts.

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10. The granular product or tablet according to claim 9, wherein the granules also contain at least one component selected from the group consisting of binders - preferably polyvinylpyrrolidone (PVP) -, small amounts of a tenside -preferably selected from the group consisting of dioctyl sodium sulfosuccinate and sodium lauryl sulfate -, alkali and/or alkaline earth carbonate and/or bicarbonate.

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11. The granular product or tablet according to any one of the preceding claims, wherein it contains, with respect to the total mixture, about 2 to about 30 % by weight of climetidine; about 30 % by weight of a solid, edible organic acid; about 12 to about 40 % by weight of at least one alkali or alkaline earth carbonate or bicarbonate (of which about 2 to about 10 % by weight is sodium carbonate as moisture-binding agent); about 1 to about 4 % by

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weight of a sweetener; about 0.01 to about 30 % by weight of a neutral substance (of which about 0.01 to about 0.05% by weight is for the neutral substance coating), preferably about 3 to about 20 % by weight of sorbitol and about 2 to about 10 % by weight of mannitor, about 0.005 to about 0.5 % by weight of an antitioanning agent, and about 0.1 to about 3 % by weight of flavoring agent.

- 12. The granular product or tablet according to any one of claims 1 to 10, wherein it contains, with respect to the total mixture, the following components: about 0.4 to about 4.5% by weight of cisapride; about 0.4 to about 4.5% by weight of suspending agent; about 0.1 to about 1.8 by weight of binder, prefeatably polyvinyprindidone (PVP); about 0.03 to about 0.35% by weight of tenside, preferably diodyl sodium sulfosuccinate; about 30 to about 55% by weight of tenside, preferably diodyl sodium sulfosuccinate; about 30 to about 55% by weight of at least one alfalia or affaitaborate of bicarborate (of which about 0.2 to about 10% by weight of at least one alfalia or alkaliarine earth ratborate or bicarborate (of which about 0.2 to about 10% by weight of neutral substance (of which about 0.22 to about 0.1% by weight is for the neutral substance coating), preferably selected from the group consisting of maltodextrin, lactose and mannitor, about 0.056 to about 0.05% by weight of antitioanming and, preferably selected from the group consisting of dimethicone, and about 0.22 to about 5.8% by weight of articoan.
- 13. The granular product or tablet according to any one of claims 1 to 10, wherein it contains, with respect to the total mixture, the following components:
- about 0.1 to about 0.5 % by weight of beta-carotene (100%);

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- about 0 to about 2 % by weight of tocopheryl acetate (100%);
 about 35 to about 70 % by weight of solid, edible organic acid, preferably about 0 to about 10 % by weight of
- ascorbic acid, about 35 to about 55 % by weight of citric acid, and about 0 to about 5 % by weight of malic acid; about 11 to about 38 % by weight of at least one alkali or alkaline earth carbonate or bicarbonate, preferably about 5 to about 15 % by weight of calcium carbonate and about 5 to about 15 % by weight of calcium carbonate and about 5 to about 20 % by weight of sodium bicar-
- about 1 to about 4 % by weight of sweetener;
- about 0.1 to about 35.0 % by weight of neutral substance (of which about 0.1 to about 0.5 % by weight is for the neutral substance coating), preferably about 1 to about 10 % by weight of soibitol and about 5 to about 25 % by weight of mannitol; and

- about 0.3 to about 3 % by weight of flavouring.
- 14. The granular product or tablet according to any one of claims 1 to 10, wherein it contains, with respect to the total mixture, the following components: about 3 to about 14 % by weight of rantitidine hydrochloride (75 300 nig per dose); about 30 to about 50 % by weight of citric acid; about 0 to about 20 % by weight of morrosodium citate; about 10 to about 30 % by weight of sodium bicarbonate; about 2 to about 10 % by weight of sodium reabonate; about 1 to about 3 % by weight of sweetener; about 0.5 to about 10 % by weight of neutral substance for the first coating as well as about 0 to about 15 % by weight of additional neutral substances; about 0 to about 8 % by weight of flavoring.
- 15. An effervescent tablet containing at least one pharmaceutically active substance and an effervescent system comprising at least one solid, edible, organic acid, at least one alkali metal carbonate or bicarbonate as a gas-forming component and at least one alkali metal salt of the acid, at least two layers being applied to carrier crystals constitution of the attest one acid, wherein the first layer contains at least one office, organic acid or the alkali metal salt of this other acid, or both, whereas the sectord layer contains at least one alkali metal salt of said at least one acid, and wherein the first layer additionally contains a neutral substance selected from the group consisting of a water-soluble polymer, a higher alcohol, a cabobydrate and a hydrocolloid.
- 50 16. A granular product or tablet with an effervescent system according to any one of claims 1 15 and clsaputde as the pharmaceutically active substance, wherein, at a total weight of less than 2 grams, preferably less than about 1.6 grams, it has an acid-binding capacity of less than 5 meq, preferably less than 3 meq.
- 17. A granular product or tablet with an effervescent system according to any one of claims 1 15 and cinnetidine as the pharmaceutically active substance, wherein, at a total weight of less than 2.5 grams, preferably less than about 2.0 grams, it has an acid-binding capacity of less than 5 meg, preferably less than 3 meg.
- A granular product or tablet with an effervescent system according to any one of claims 1 15 and rantItIdIne as

the pharmaceutically active substance, wherein, at a total weight of less than 2.6 grams, preferably less than 2.0 g, it has an acid-binding capacity of less than 3 meq, preferably less than 2 meq. 19. A method for the preparation of a granular product or of a tablet according to any one of the preceding claims, wherein crystals of at least one solid, edible organic acid are wetted with an aqueous solution of a neutral substance, and then, before complete of ying, an alkali and/or alkaline earth carbonate and/or bicarbonate in powder farm is unitomly distributed and anchored on the moist surface layer by mixing, whereupon the effer-vescent grains thus prepared are dried and mixed with a pharmaceutically acive substance - preferably with an acid-sensitive one, especially one that is selected from the group consisting of H2-blockers, climetidine, rantifidine, clisapride and beta-carolene - and pharmaceutically acceptable adjuvants, and optionally compressed into tablets.

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- 20. The method according to claim 19, wherein, on the effervescent grains, at least one additional coating is applied by wetting the grains with the solution of a buffer substance, preferably one that is selected from the group consisting of alketif carbonate, alketif bicarbonate, alketif bicarbonate, alketif bicarbonate, alketif bicarbonate, alketif and a solid edible organic acid and alketime earth satil or at least one solid edible organic acid and alketime earth satil or at least one solid edible organic acid.
- 21. The method according to claim 19 or 20, wherein the solution further comprises a neutral substance selected from the group consisting of a water-soluble polymer, a higher alcohol, a carbohydrate and a hydrocolloid.
- 22. The meltind according to any one of claims 19 to 21, wherein, in addition to the drug, the effervescent grains are also mixed with a granular product which has been made by applying an antifoaming agent in an appropriate solvent to the surface of neutral substance particles, and drying the solvent.
- 23. The method according to any one of claims 19 to 22, wherein the dried effervescent grains are wetted with ethanol, which preferably contains an antifoaming agent dissolved, and are dried again, by evaporating the ethanol, to remove residual moisture.
- 24. The method according to any one of claims 19 to 23, wherein the pharmaceutically active substance, before admixing it to the effervescent system, is together with a binding agent and/or a tenside applied in solution to and uniformly distributed on the grains of a suspension agent and dried.

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25. The method according to any one of claims 19 to 24, wherein the pharmaceutically active substance, before admixing it to the effervescent system, is mixed with at least one neutral substance, at least one suspension agent and at least one substance selected from the group of alkali carbonate, alkali bicarbonate, alkaline earth carbonate, alkaline earth bicarbonate, alkaline at the size of a size

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- 26. A process for the manufacture of effervescent granules from a powdered or granular mixture of a solid, eclible, organic acid and the carbonate and/or bicarbonate of an alkali and/or alkaline earth metal under vacuum, wherein for the passivation of the surface of at least one of the components to a state of stong inertia to the reaction, there is added to the heated mixture during the treatment under vacuum a melered quantity of a polar solvent, the difference in pressure caused by development of carbon dioxide through the addition of solven during the reaction being determined up to a maximum of 1000 mbar, the volume and mass of the carbon dioxide iberated being ascertained from this difference in pressure, and the heat treatment being repeated, after rapid drying of the mixture, as many times as necessary to obtain passivation of the surface as indicated by an evident slowing down of the reaction and by a reduced development of gas, and wherein in said polar solvent there is dissolved a neutral substance selected from the group consisting of a water-solvible polymer, a higher alcohol, a carbohydrate and a hydrocollicit.
- 27. A process for the preparation of an effervescent granular material containing at least one solid, crystalline edible organic acid and at least one carbonate of an alkali metal or an alkaline earth metal which splits off CO₂ upon reaction with said organic acid in aqueous solution, which comprises:
- pre-reacting a portion of said organic acid and said carbonate in solution in water and/or alcohol to form a prereaction product

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adding said pre-reaction product to an additional portion of said organic acid in crystalline form with thorough mixing to form a first coating by reaction with said organic acid crystals and liberation of the resulting water of crystallization,

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applying at least one additional coating including said carbonate onto the organic acid crystals with said first coating adhering thereto, and

lerminating the reaction after the last coating has been applied by drying, wherein to said pre-reaction product there is added a neutral substance selected from the group consisting of a water-soluble polymer, a higher alcohol, a carbohydrate and a hydrocolloid.

Patentansprüche 4 4 1

Ein granuläres Brauseprodukt, welches zum Herstellen einer wässrigen Lösung oder Suspension einer oder mohrere pharmazeulisch aktiver Substanzen zur oraten Verabreichung geeignet ist, und welches in Tabhelten preestbar ist, und/oder dieses Produkt in Tabheltenform, uni Brausekönnern, die von Tagerkristallen wenigslens einer lesten, genieszbaren organischen Saure erhalten worden sind, welche im wesentlichen mit wenigslens einer lesten, genieszbaren organischen Saure erhalten worden sind, welche im wesentlichen mit wenigslens einer Beschichtung bedeckt sind, die mindestens eine wassenfosliche, neutrale Substanz erntrält, wobei die neutrale Substanz zum Absenken des Schmelzpunktes der Sauredvistalle an ihrer Oberläche wirksam ist, und wenigstens eine aus der aus Alkalicatonat, Alkalibicatonat, Erdalkalicatonat, einem Alkalisaz, und einem Erdalkalisalz zum indest einer festen, gereiessbaren organischen Saure bestehenden Gruppe wenigstens einer festen, geneiessbaren organischen Saure beschichung angebacht ist.

 Granuláres Produkt oder Tablette nach Anspruch 1, wobei die neutrale Substanz aus der aus einem wassenlöslizo chen Polymer einem höheren Alkhold, einem Köhlehydrat und einem Hydrokolloid bestehenden Gruppe ausgewaht ist, welche neutrale Substanz in einer Menge von etwa 0,05 bis annahernd 1,0 Gewichts-%, vorzugsweise von etwa 0,07 bis ungelähr 0,8 Gewührs-%, vorhanden ist. Granulares Produkt oder Tablette nach Anspruch 1 oder 2, wobei ein Feuchtigkeltsblindernittel an den Brausekörnern verankert ist, welches Feuchtigkeitsbindemittel vorzugsweise aus der aus katziniertem Soda und Natirumsulfat bestehenden Gruppe ausgewählt ist und vorzugsweise in einer Menge von etwa 4 bis ungelähr 10 Gewichts%, bezogen auf die gesamte Mischung, eingesetzt ist.

Granulares Produkt oder Tablette nach einem der vonhergehenden Ansprüche, wobei wenigstens eine zusätzliche
Beschlichtung an den Brausekörnern angebracht ist, welche eine aus der aus Alkalisatzen und/oder Erdalkalisatzen wenigstens einer Testen, geniessbaren organischen Saure bestehenden Gruppe ausgewählte Substanz als
Pulfer und gegebenenfalls eine zusätzliche neutrale Substanz aufweist, und wobei vorzugsweise wenigstens eine
der Beschichtungen ein Antischaummittel enthält.

 Granuläres Produkt oder Tablette nach einem der vorhergehenden Ansprüche, wobei das granuläre Produkt oder das in Tablettenform gepresste granuläre Produkt ferner mindestens ein in einem eigenen granulären Produkt vorhandenes Antischaummittel autweist. Granuläres Produkt oder Tablette nach Anspruch 4 oder 5, wobei das Antischaummittel aus der aus Dinnethioon und Sinnethicon bestehenden Gruppe ausgewählt ist und in einer Menge von etwa 0,005 bis ungelähr 0,5 Gewichts-%, bezogen auf die gesamte Mischung, oder von etwa 0,05 bis ungelähr 2,0 Gewichts-%, bezogen auf die pharmazautisch aktive Substanz, eingesetzt ist. Granuläres Produkt oder Tablette nach einem der vorhergehenden Ansprüche, wobei es bzw. sie eine Säureblindungsfähligkelt von weniger als 5, vorzugsweise weniger als 3 meg, gemessen nach USP XXII, aufweist.

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Granulaires Produkt oder Tablette nach einem der vorhergehenden Ansprüche, wobei es bzw. sie bei einem Gesamtgewicht von nicht mehr als 2.5, vorzugsweise nicht mehr als 2.0 Gramm in Wasser bei Rauntemperatur eine Autlösungszelt von weniger als 180, vorzugsweise weniger als 120, Sekunden aufweist. Granulares Produkt oder Tablette nach einem der vorhergehenden Arsprüche, mit einer hydrophoben planmazeutisch aktiven Substanz, wobei die hydrophobe Substanz in von den Brausekomponenten gesonderten Granula
vorliegt, in welchen Granula die hydrophobe Substanz auf wenigstens einer aus der aus Suspendiermitteln - welche vorzugsweise aus der aus Aerosil[®] und Avicel[®] bestehenden Gruppe gewählt sind - und neutralen Substanse - welche vorzugsweise aus der aus Mannich und Sorbitol bestehenden Gruppe gewählt sind - bestehenden
Gruppe ausgewähllen Substanz geschrichel bzw. an Ilnnen verankent ist.

. Granulares Produkt oder Tablette nach Anspruch 9, wobei die Granula auch wenigstens eine aus der aus BIndern

 vorzugsweise Polyvinylpyrrolidon (PVP) ·, geringen Mengen eines Tensids · welches vorzugsweise aus der aus Dioctyl-Natriumsulfosuccinat und Natriumlaurylsulfat bestehenden Gruppe gewählt ist ·, Alkali- und/oder Erdalkalicarbonat und/oder -bicarbonat bestehenden Gruppe ausgewählte Komponente enthalten. 11. Granulares Produkt oder Tablette nach einem der vorhergehenden Ansprüche, wobei es bzw. sie, bezogen auf die gesamte Mischung, etwa 20 bis ungefahr 80 Gewichts-% Cinnetidin, etwa 30 bis ungefahr 60 Gewichts-% einer festen, geniessbaren rognischen Saure, etwa 12 bis ungefahr 40 Gewichts-% wenigstens eines Aklai- oder Erdalkalicrationals oder -bicarbonats (wovon etwa 2 bis ungelahr 10 Gewichts-% Institumcarbonat als Feuchtigheits-bindernitlel ist); etwa 1 bis ungefahr 4 Gewichts-% eines Süssstolfes, etwa 0,01 bis ungefahr 30 Gewichts-% einer neutralen Substanz (wovon etwa 0,01 bis ungefahr 0,05 Gewichts-% für die Beschichtung mit neutraler Substanz ist), vorzugsweise etwa 3 bis ungefahr 20 Gewichts-% since dewa 2 bis ungefahr 3 Gewichts-% eines Geschnadoknitiels.

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12. Granuláres Produkt oder Tablette nach einem der Ansprüche 1 bis 10, wobei es bzw. sie, bezogen auf die gesamte Mischung, die folgenden Komponenten enthält: etwa 0,4 bis ungefähr 4,5 Gewichts-% Eines Suspendiermitlets; etwa 0,1 bis ungefähr 1 Gewichts-% Einer, vorzugsweise Polyvinry-pyrolidon (PVP): etwa 0,03 bis ungefähr 0,35 Gewichts-% Prensid, vorzugsweise Brotyringt-pyrolidon (PVP): etwa 0,03 bis ungefähr 5,35 Gewichts-% einer festen, geniessbaren organischen Saure, vorzugsweise Zithonensäure; etwa 12 bis ungefähr 50 Gewichts-% einer festen, geniessbaren organischen Saure, vorzugsweise Zithonensäure; etwa 12 bis ungefähr 10 Gewichts-% einer Raten, unfoder Erdakfalicarbonats oder - bicanbonats (wovon etwa 2 bis ungefähr 10 Gewichts-% Natriumcarbonat als Feuchtigkeitsbindemittel sind); etwa 0,3 bis ungemen auf 0,02 bis ungefähr 0,1 Gewichts-% eines Abstanz (wovon etwa 0,02 bis ungefähr 0,1 Gewichts-% eines Abstanz ist), die vorzugsweise aus der aus Mallodextrin, Laktose und Mannitol bestehenden Gruppe ausgewählt ist; etwa 0,03 bis ungefähr 0,05 Gewichts-% eines Antischaummittels, welches vorzugsweise aus der aus Dimethicon und Simethicon bestehenden Gruppe ausgewählt ist, und etwa 0,2 bis ungefähr 5 Gewichts-% eines Geschmadsernitiels.

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- Granulares Produkt oder Tablette nach einem der Ansprüche 1 bis 10, wobei es bzw. sie, bezogen auf die gesamte Mischung, die folgenden Komponenten enthält:
- etwa 0,1 bis ungefähr 0,5 Gewichts-% beta-Carotin (100%);

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- etwa 0 bis ungefähr 2 Gewichts-% Tocopherylazetat (100%);
- etwa 35 bis ungelfähr 70 Gewichts. % einer festen, geniessbaren organischen Säure, vorzugsweise etwa 0 bis ungefähr 10 Gewichts. % Ascotbinsäure, etwa 35 bis ungefähr 55 Gewichts. % Zitronensäure und etwa 0 bis ungefähr 5 Gewichts. % Maleinsäure;
- etwa 11 bis ungetâtv 38 Gewichts-% wenigstens eines Alkali- oder Erdalkalicarbonats oder -bicarbonats, vorzugsweise etwa 5 bis ungefâtv 15 Gewichts-% Calciumcarbonat und etwa 5 bis ungelâtv 20 Gewichts-% Natriumbicarbonat;
- elwa 1 bis ungefähr 4 Gewichts-% eines Süssstoffes;

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- etwa 0,1 bis ungefähr 35,0 Gewichts-% einer neutralen Substanz (wovon etwa 0,1 bis ungefähr 0,5 Gewichts% für die Beschichtung mit neutraler Substanz ist), die vorzugsweise etwa 1 bis ungefähr 10 Gewichts-% Sorbitol und etwa 5 bis ungefähr 25 Gewichts-% Mannitol sind; und
 - etwa 0,3 bis ungefähr 3 Gewichts-% eines Geschmacksmittels.
- 14. Gaauulares Produkt oder Tablette nach einem der Ansprüche 1 bis 10, wobel es bzw. sie, bezogen auf die gesamte Mischung, die Digenden Komponenten enthält: etwa 3 bis ungelähr 14 Gewichts.* Ranitidin-Hydrochlorid (75-300 mg pro Dosis), etwa 30 bis ungelähr 30 Gewichts.* Zitronersäure; etwa 0 bis ungelähr 20 Gewichts.* Zitronersäure; etwa 0 bis ungelähr 20 Gewichts.* Natriumbicarbonat; etwa 2 bis ungelähr 10 Gewichts.* Natriumbicarbonat; etwa 10 bis ungelähr 3 Gewichts.* Natriumbicarbonat; etwa 0.05 bis ungelähr 3 Gewichts.* Natriumbicarbonat; etwa 0.05 bis ungelähr 10 Gewichts.* einer neturäeln Substanz für die erste Beschichtung sowie etwa 0 bis ungelähr 15 Gewichts.* zusätzlicher neturäer Substanzzen; etwa 0.1 bis ungelähr 4 Gewichts.* eines Geschmadsrantiels.
- 15. Eine Brausetablette, welche wenigstens eine pharmazeutisch aktive Substianz und ein Brausesystem mit wenigstens einer Brausesystem mit wenigstens einer Besten, geniessbaten organischen Säure, wenigstens einem Alkalimetalicarbonat oder blicarbonat als gasbildende Kontponente und mindestens einem Alkalimetalistat der Säure, wobei zumindest zwei Schlichten auf Tägerkristalle aufgettagen sind, welche aus der wenigstens einen Säure bestehen, wobei die erste Schicht zumindest eine weitere feste, gemiesbate organische Säure oder das Alkalimetalistat dieser weiteren Säure oder

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beide enthält, wogegen die zweite Schicht mindestens ein Alkalimetallsalz der wenigstens einen Säune enthält, und wobei die erste Schicht zusätzlich eine aus der aus einem wasserlöslichen Polynner, einem höheren Alkohol, einem Kohlerydrat und einem Hydrocolloid bestehenden Gruppe ausgewählte neutrale Substanz enthält.

- 6 16. Granuläres Produkt oder Tablette mit einem Brausesystem nach einem der Ansprüche 1 bis 15 und Cisaprid als pharmazeutisch aktive Substanz, wobei, bei einem Gesamigewicht von wentger als 2 Gannn, vorzugsweise weniger als elwa 1,6 Gramm, es bzw. sie eine Saurebindungsfähigkeit von wentger als 5 meg, vorzugsweise wentger als 3 meg, besitzt.
- 17. Granuläres Produkt oder Tablette mit einem Brausesystem nach einem der Ansprüche 1 bis 15 und Ginetidin als pharmazeutisch aktive Substanz, wobei, bei einem Gesamtigewicht von wentger als 2,5 Gramm, vorzugsweise wentger als ehwa 2.0 Gramm, es bzw. sie eine Saurebindungsfähigkeit von wentger als 5 nieg, vorzugsweise weniger als 3 meg, besitzt.
- 16 Granuläres Produkt oder Tablette mit einem Brausesystem nach einem der Ansprüche 1 bis 15 und Rantilldin als pharmazeutisch aktive Substanz, wobei, bei einem Gesamtgewicht von weniger als 2,6 Grannu, vorzugsweise weniger als 2,0 Grannu, es bzw. sie eine Saurebindungsfähigkeit von weniger als 3 meg, vorzugsweise weniger als 2 meg, besitzt.
- Verfahren zur Herstellung eines granuflaren Produktes oder einer Tablette nach einem der vorhergehenden Ansprüche, bei dem Kristalle wenigstens einer festen, geniessbaren organischen Säure mit einer wässrigen I. üsung einer neutralen Süustanz angeleuchtet wird und dann vor dem vollständigen Trocknen ein Aktali- und/oder Erdalkalicarbonat und/oder Erdalkalicarbonat und/oder Erdalkalicarbonat und/oder Erdalkalicarbonat in Pulverform gelichmässig verteilt und an der feuchten Oberflächenschicht durch Mischen verankert wird, worauf die so hergestellten Brausekkriner getrochent und mit einer priarmazeutisch aktiven Substanz vorzugsweise mit einer säurenerpfündlichen, insbesondere einer aus der aus H2-Blockern, Gimetidin, Ramitidin, Cisaprid und beta-Carotin bestehenden Gruppe ausgewählten und phairmazeutisch aktzeptablen Hilsmittlein genrischt, und gegebenenfalls zu Tabletten gepresst, werden.
- 20. Verfahren nach Anspruch 19, bei dem auf den Brausekörnern mindestens eine zusätzliche Beschlehtung durch so Beleuchten der Körner mit der Lösung einer Puffersubstanz aufgebracht wird, vorzugsweise einer solchen, welche aus der aus Alkalicarbonat, Erdalkalicarbonat, Erdalkalicarbonat, einem Alkalisatz zumindest einer festen, geniessbaren organischen Saure und einem Erdalkalisatz zumindest einer festen, geniessbaren organischen Saure bestehenden Gruppe ausgewählt ist.
- 21. Verfahren nach Anspruch 19 oder 20, bei dem die Lösung ferner eine aus der aus einen wassenlöslichen Polymer, einem h
 öheren Alkohol, einem Kohlehydrat und einen Hydrocolloid bestehenden Gruppe ausgew
 ältli ist.
- 22. Verfahren nach einem der Ansprüche 19 bis 21, bei dem, zusätzlich zum Arzneimittel, die Brauset/drner auch mit einem granulären Produkt gemischt werden, das durch Auftragen eines Antischaummittels in einer geeigneten Lösung auf die Obertläche von Partikeln einer neutralen Substanz hergestellt worden ist, und das Lösungsmittel getrodnet wird.

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23. Verfahren nach einem der Ansprüche 19 bis 22, bei dem die getrockneten Brausekörner mit Athanol befeuchtet, das vorzugsweise ein Antischaummittel gelöst enthält, und durch Verdampfen des Äthanols wieder getrocknet werden, um die Restleuchtigkeil zu beseitigen.

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- 24. Verfahren nach einem der Ansprüche 19 bis 23, bei dem die pharmazeutisch aktive Substanz, vor ihrer Zumischung zum Brausesystem, in Lösung zusammen mit einem Bindemittel und/oder einem Tensid auf die Könner eines Suspendiermittels aufgetragen und gleichmassig verteilt und getrocknet wird.
- 25. Verfahren nach einem der Ansprüche 19 bis 24, bei dem die pharmazeutisch aktive Substanz, vor ihrer Zumitschung zum Brausesystem, mit wenigstens einer neutralen Substanz, mindestens einem Suspendiermittel und zunindest einer aus Aleafachonat, Alkalbioarbonat, Endakalizabonat, Endakalisibarbonat, einem Alkalisaliz zumindest einer desten, genessbaren organischen Säure und einem Erdalkalisalz zumindest einer festen, geniessbaren organischen Säure und einem Erdalkalisalz zumindest einer festen, geniessbaren organischen Saure bestehenden Gruppe ausgewählten Substanz gemischt wird, worauf die Lösung wenigsteins eines Britchmittels unz/order eines Tensids zumindest einmat auf die Könner der Mischung aufgetlagen, wentalt und nach der Mischung aufgetlagen.

26. Verfahren zur Herstellung von Brausegranula aus einer pulverfürmigen oder granufaren Mischung einer festen, geniessbaren organischen Saure und dem Cabonat undoder Braatbonat eines Afkalt- undoder Erdalkalimetalis unter Vakuum, bei dem zur Passivierung der Oberfläche wertigstens einer der Komponenten zu einem Zustand siahker Tragheit gegenüber der Readkoin der erhitzten Mischung wahrend der Behandung unter Vakuum eine dosierte Menge eines polaren Lösungsmittels zugefügt wird, die durch die Entwickfung von Kohlendfoxyd durch die Zugabe des Lösungsmittels während der Reaktion verursachte Druckdrifterenz bis auf ein Maximum von 1000 bar bestimmt wird, wobei das Volumen und die Masse des freigesetzert follbendfoxyds aus dieser Druckfilferenz bis motwer einnittell wird, und die Warmebehandfung nacht aschem Trochen der Mischung so oft wiedenholt wird, als notwerdig ist, um die Passivierung der Oberfläche zu erhalten, wie durch eine deutliche Verlangsamung der Reaktion und eine verningste Passenwickung angezeigt wird, und wobei in der polaren Lösung eine aus einem wasserlöstlichen Polymer, einem Hoheren Alkohol, einem Kohlehydrat und einem Hydrocolloid bestehenden Gruppe ausgewählte neutrale Suskalanz gelost wird, einem Rohlehydrat und einem Hydrocolloid bestehenden Gruppe ausgewählte neutrale Suskalanz gelost wird.

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27. Verfahren zur Herstellung von granulärem Brausematerial, welches mindestens eine feste, geniessbare organische Saure und zumindest ein Carbonat eines Alkali- oder Erdalkalimetalls enthält, das bei Reaktion mit der organischen Saure in einer wässrigen Lösung CO₂ abgibt, welches folgendes aufweist:

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- vorab Unsetzen eines Teiles der organischen Säure und des Carbonats in einer Lösung in Wasser und/oder Alkohol, um ein Vorreaktionsprodukt zu schaffen,
- Zugeben des Vorreaktionsproduktes zu einem weiteren Teil der organischen Saure in kristalliner Form unter sorgtaltigem Mischen, um durch Reaktion mit den Kristallen der organischen Saure und der sich daraus ergebenden Freisetzung von Kristallisationswasser eine erste Beschichtung zu bilden,

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- Aufbringen wenigstens einer weiteren, das Carbonat aufweisenden Beschichtung auf den Kristallen der organischen Säure, an denen die erste Beschichtung anhaftet, und
- Abschliessen der Reaktion, nachdem die latzte Beschichtung aufgetragen worden ist, durch Trocknen, wobei eine aus der aus einem wasserlöslichen Polymer, einem höheren Alkohol, einem Köhlehydrat und einem Hydiocolloid bestehenden Gruppe ausgewählte neutrale Substanz dem Vorreaktionsprodukt hinzugelügt wird.

Revendications

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1. Produit effervescent granulé, convenant pour la préparation d'une suspension ou d'une solution aqueuse d'une substance active du point de vue pharmaceutique ou davanlage, destiné à une administration orale, et susceptible d'être pressé en comprimés buou ledit produit sous forme de comprimés, comprenant des grains effervescents obtenus à partir de cristaux porteurs d'au moins un acide organique alimentaire et solide, qui sont sensiblement recouverts par au moins un revêtement connemant au moins un excelement connemant au moins une substance neutre hydrosoluble, dans leque ladite substance neutre est capable d'abaisser le point de fusion des cristaux d'acide à leur surface, et au moins une substance - choisie dans le groupe constitué par les carbonates alcalins, les bicarbonates alcalins des carbonates alcalins-lerreux d'au moins un acide organique alimentaire et solide - est appliquée sur ledit revêtement.

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 Produit granulé ou comprimé selon la revendication 1, dans lequel la substance neutre est choisie dans le groupe constitué par les polymères hydrosolubles, les alcools supérieurs, les hydrates de carbone et les hydrocolloïdes et dans lequel fadite substance neutre est présente en une quantité allant d'environ 0,05 à environ 1,0 % en poids et, de préférence, d'environ 0,07 à environ 0,8 % en poids.

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- Produit granulé ou comprimé selon la revendication 1 ou la revendication 2, dans lequel un agent fixant l'humidité
 est fixé sur lesdits grains effervescents, cet agent fixant l'humidité étant choisi, de préférence, dans le groupe constitué par le carbonate de sodium anhydre et le sulfate de sodium anhydre et étant appliqué, de préférence, en une
 quantité allant d'environ 4 à environ 10 % en poids, par rapport au mélange total.
- 4. Produit granudé ou compnimé selon l'une queliconque des revendications précédentes, dans lequel on a appliqué sur les grains effervescents au moins un revêtement additionnel, comprenant une substance choisie dans le groupe constitué par les sels alcalins eviou les sels alcalins eviou les sels alcalins eviou les sels alcalino-terreux d'au moins un acide organique alimentaire et solide servant de lampon et, à titre facultait, une substance neutre additionnelle et dans lequel, de préférence, au moins un des révêtements contient un appet antimousse.

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Produit effervescent granulé ou comprimé selon l'une quelconque des revendications précédentes, dans lequel le

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produit granulé ou ledit produit granulé pressé sous forme de comprimés comprend, en outre, au moins un agent antimousse présent lui-même sous forme d'un produit granulé séparé.

- 6. Produit granulé ou comprimé selon la revendication 4 ou la revendication 5, dans lequel l'agent antimousse est choisi dans le groupe constituté par la dimethicone et la siméthicone et est applique en une quantilé d'environ 0,005 à environ 0,005 par rapport à la substance active du point de vue pharmeceutique.
- Produit granulé ou comprimé selon l'une quelconque des revendications précédentes, ayant une capacité de fixation d'acides inférieure à 5 et, de préférence, inférieure à 3 méq., la détermination étant faite selon USP XXII.
- 8. Produit granulé ou comprimé selon l'une quelconque des revendications précédentes, qui, pour un poids total ne dépassant pas 2,5 et, de préférence 2,0 grammes, présente un temps de dissolution dans l'eau à la température ambiante intérieur à 180 et, de préférence, intérieur à 120 secondes.
- 9. Produit granulé ou comprimé selon l'une quelconque des revendications précédentes, qui comprend une substance active du point de vue pharmaceulique et hydrophobe, et dans lequel la substance hydrophobe est présente dans des granules distincts des composants effervescents, la substance hydrophobe de ces granules étant appliquée en revêtement ou fixée sur au moins une substance choisie dans le groupe constitué par des agents de sus pension (choisies, de préférence, dans le groupe constitué par le produit Aerosi ® et le produit Avricel ®) et des substances neutres (choisies, de préférence, dans le groupe constitué par le mannitot et le produit Avricel ®) et des substances neutres (choisies, de préférence, dans le groupe constitué par le mannitot et le sorbitol).
- 10. Produit granulé ou comprimé selon la revendication 9, dans lequel les granules contiennent également au moins un composant choisi dans le groupe constitué par des liants (de préférence la polyvinythynolidone (PVP)), de petites quantités d'un tensicacití (choisi, de préférence, dans le groupe constitué par le dioctyl-sulfosuccinate de sodium et le lauryl-sulfate de sodium), et les carbonates et/ou les bicarbonates alcalins et/ou alcalino-terreux.
- 11. Procluit granulé ou comprimé selon l'une quelconque des revendications précédentes contenant, par rapport au melange foital, d'environ 2 à environ 30 % en poids de cimiéticine; d'environ 60 % en poids d'un acide organique alimentaire et solide; d'environ 10 % en poids d'un moins un catbonate en un bicarbonate alcain ou alcalimoterreux (dont d'enviror 12 à environ 10 % en poids d'au moins un catbonate et socialimoterreux (dont d'enviror 2 à environ 10 % en poids cent constitués par le carbonate de socialimoterreux (dont d'enviror 1 à environ 4 % en poids cent constitués par le carbonate de socialiment. Elsé comme agent fixant l'humidité); d'environ 1 à environ 1 % en poids sur edulcorant; d'environ 0,01 à environ 30 % en poids d'une substance neutre) (dont d'environ 0,01 à environ 0,05 % en poids de carbonate de d'environ 0,05 à environ 0,05 % en poids d'un agent antimousse et d'environ 0,1 à environ 3 % en poids de marnitot; d'environ 0,005 à environ 0,5 % en poids d'un agent antimousse et d'environ 0,1 à environ 3 % en poids d'un agent antimousse et d'environ 0,1 à environ 3 % en poids d'un agent antimousse et d'environ 0,1 à environ 3 % en poids d'un agent antimousse et d'environ 0,1 à environ 3 % en poids d'un agent antimousse et d'environ 0,1 à environ 3 % en poids d'un agent antimousse et d'environ 0,1 à environ 3 % en poids d'un agent anonatisant.
- 12. Produit granulé ou comprimé selon l'une quelconque des revendications 1 à 10, contenant, par rapport au melanige total, les composants suivants : d'environ 0,4 à environ 4,5 % en poids de cisapride; d'environ 0,4 à environ 4,5 % en poids d'un agent de suspension; d'environ 0,1 à environ 1 % en poids d'un liant, de préférence la polyvinylpyrroldione (PVP); d'environ 0,03 à environ 0,35 % en poids d'un tensioaciti, de préférence la polyvinylpyrroldione (PVP); d'environ 30 à environ 55 % en poids d'un moins un carbonate ou un bicarbonate alcalin ou alcalin-quer environ 12 à environ 40 % en poids d'un moins un carbonate ou un bicarbonate alcalin ou alcalin-quer eux (dont d'environ 2 à environ 0,0 % en poids sont constitués par le carbonate de sodium utilisé comme agent fixant l'humictie); d'environ 0,3 à environ 2,5 % en poids d'un édulcorant; d'environ 0,02 à environ 0,0 % en poids envien poids servent pour le revêtement de la substance neutre), choisie, de préférence, dans le groupe constitué par la mallodatrine, le lactose et le mannitot; d'environ 0,005 % en poids d'agent anilimousse, choisi, de préférence, dans le groupe constitué par la mallodatrine, le lactose et le mannitot; d'environ 0,005 % en poids d'agent anilimousse, choisi, de préférence, dans le groupe constitué par la mallodatrine, le lactose et le mannitot d'environ 0,005 % en poids d'agent anilimousse, choisi, de préférence, dans le groupe constitué par la dimétrioone et la sindétricone, a d'environ 0,2 à environ 5 % en poids d'un agent aonnatisant.

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- Produit granulé ou comprimé selon l'une quelconque des revendications 1 à 10, contenant, par rapport au mélange total. les composants suivants:
- d'environ 0,1 à environ 0,5 % en poids de bêta-carotène (100 %);
- d'environ 0 à environ 2 % en poids d'acétate de tocophérol (100 %),

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d'environ 35 à environ 70 % en poids d'un acide organique alimentaire et solide, de préférence d'environ 0 à
environ 10 % en poids d'acide ascolbique, d'environ 35 à environ 55 % en poids d'acide citrique et d'environ 0
à environ 5 % en poids d'acide malique;

- d'environ 11 à environ 38 % en poids d'au moins un carbonate ou un bicarbonate alcalin ou alcalino-terreux, de préférence d'environ 5 à environ 15 % en poids de carbonate de calcium et d'environ 5 à environ 20 % en poids de bicarbonate de sodium;
- d'environ 1 à environ 4 % en poids d'un édulcorant;
- d'environ 0,1 à environ 35,0 % en poids d'une substance neutre (dont d'environ 0,1 à environ 0,5 % en poids servent pour le revêtement de la substance neutre), de préférence d'environ 1 à environ 10 % en poids de sorbitol et d'environ 5 à environ 25 % en poids de mannitol; et
- d'environ 0,3 à environ 3 % en poids d'un agent aromatisant
- total, les composants suivants : d'environ 3 à environ 14 % en poids de chlorhydrate de ranitidine (75 300 mg par tances neutres additionnelles; d'environ 0 à environ 8 % en poids de granules d'un agent antimousse et d'environ Produit granulé ou comprimé selon l'une quelconque des revendications 1 à 10, contenant, par rapport au mélange dose); d'environ 30 à environ 50 % en poids d'acide citrique; d'environ 0 à environ 20 % en poids de citrate monosodique; d'environ 10 à environ 30 % en poids de bicarbonate de sodium; d'environ 2 à environ 10 % en poids de carbonate de sodium; d'environ 1 à environ 3 % en poids d'un édulcorant; d'environ 0,05 à environ 0,2 % en poids de la substance neutre utilisée pour le premier revêtement ainsi que d'environ 0 à environ 15 % en poids de subs-0,1 à environ 4 % en poids d'un agent aromalisant. 4. 2 5
- vescent comprenant au moins un acide organique alimentaire et solide, au moins un carbonate ou un bicarbonate de métal alcalin en tant que composant générant du gaz et au moins un sel de métal alcalin de l'acide, dans lequel au moins deux couches sont appliquées aux cristaux porteurs constitués par au moins un premier acide, la première couche contenant au moins un autre acide organique alimentaire et solide ou un sel de métal alcalin de cet la première couche contenant en plus une substance neutre choisie dans le groupe constitué par les polymères Comprimé effervescent contenant au moins une substance active sur le plan pharmaceutique et un système efferautre acide ou les deux, alors que la seconde couche contient au moins un sel de métal alcalin dudit premier acide, hydrosolubles, les alcools supérieurs, les hydrates de carbone et les hydrocolloïdes. S S
- Produit granulé ou comprimé avec un système effervescent selon l'une quelconque des revendications 1 15 et du cisapride en tant que substance active du point de vue pharmaceutique, qui, pour un poids total de moins de 2 grammes et, de préférence, de moins d'environ 1,6 grammes, a une capacité de lixation d'acides inférieure à 5 méq. et, de préférence, inférieure à 3 méq. 9 33
- Procluit granulé ou comprimé avec un système effervescent selon l'une quelconque des revendications 1 15 et de la cimétidine en tant que substance active du point de vue pharmaceutique, qui, pour un poids total de moins de 2,5 grammes et, de préférence, de moins d'environ 2,0 grammes, a une capacité de fixation d'acides inférieure à 5 méq. et, de préférence, inférieure à 3 méq 35
- la ranitidine en tant que substance active du point de vue pharmaceutique, qui, pour un poids total de moins de 2,6 Produit granulé ou conprimé avec un système effervescent selon l'une quelconque des revendications 1 - 15 et de grammes et, de préférence, de moins de 2,0 grammes, a une capacité de lixation d'acides inférieure à 3 méq. et, de préférence, inférieure à 2 méq. œ

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- Procédé de préparation d'un produit granulé ou un comprimé selon l'une quelconque des revendications précédentes, dans lequel des cristaux d'au moins un acide organique alimentaire et solide sont mouitlés avec une solution aqueuse d'une substance neutre et ensuite, avant le séchage complet, un carbonate et/ou un bicarbonate alcalin eVou alcalino-terreux, sous forme de poudre, est réparti de manière uniforme et fixé à la couche de surface humide par mélange, suite à quoi les grains effervescents ainsi préparés sont séchés et mélangés avec une substance active du point de vue pharmaceutique - qui est, de préférence, une substance sensible aux acides et qui est choisie, en particulier, dans le groupe constitué par les antagonistes des récepteurs H2, la cimétidine, la ranitidine, le cisapride et le bêta-carotène - et avec des adjuvants acceptables du point de vue pharmaceutique, puis éventuellement pressés en comprimés. 5
- additionnel, en mouillant les grains avec une solution d'une substance tampon, de préférence choisie dans le 20. Procédé selon la revendication 19, dans lequel on applique sur les grains effervescents au moins un revêtement groupe constitué par les carbonates alcalins, les bicarbonates alcalins, les carbonates alcalino-terreux, les bicarbonates alcalino-terreux, les sels alcalins d'au moins un acide organique alimentaire et solide et les sels alcalinoterreux d'au moins un acide organique alimentaire et solide. 55

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- Procédé selon la revendication 19 ou la revendication 20, dans lequel la solution contrend, en outre, une substance neutre choisie dans le groupe constitué par les polymères hydrosolubles, les alcools supérieurs, les hydrates
- vescents sont également mélangés avec un produit granulé qui a été obtenu en appliquant un agent antimousse Procédé selon l'une quelconque des revendications 19 à 21, dans lequel, en plus du médicament, les grains efferdans un solvant approprié sur la surface des particules de la substance neutre et en séchant le solvant.
- mouillés avec de l'éthanol qui contient, de préférence, un agent antimousse dissout, puis séchés à nouveau, en Procédé selon l'une quelconque des revendications 19 à 22, dans lequel les grains effervescents séchés sont évaporant l'éthanol, pour enlever l'humidité résiduelle. 23 5
- 24. Procédé selon l'une quelconque des revendications 19 à 23, dans lequel, avant de métanger la substance active du point de vue pharmaceulique au système effervescent, elle est appliquée en solution avec un agent liant et/ou un tensioactif, et répartie de manière uniforme sur les grains d'un agent de suspension et séchée. 15
- bonates alcalins, les bicarbonates alcalins, les carbonates alcalino-terreux, les bicarbonates alcalino-terreux, les Procédé selon l'une quelconque des revendications 19 à 24, dans lequel, avant de mélanger la substance active du point de vue pharmaceutique avec le système effervescent, elle est mélangée avec au moins une substance neutre, au moins un agent de suspension et au moins une substance choisie dans le groupe conprenant les carsels alcalins d'au moins un acide organique alimentaire et solide et les sels alcalino-terreux d'au moins un acide organique alimentaire et solide, suite à quoi une solution d'au moins un agent liant el/ou d'un tensioactif est appliquée et répartie sur les grains du mélange, qui sont alors séchés. 25. 8
- Procédé de fabrication de granules effervescents à partir d'un mélange pulvérulent ou d'un mélange granulé d'un de haute inertie à la réaction, on ajoute au mélange chauffé durant le traitement sous vide, une quantité mesurée d'un solvant polaire, la différence de pression provoquée par la formation de gaz carbonique produit par l'addition du solvant durant la réaction étant choisie pour atteindre au maximum 1000 mbars, le volume et la masse du gaz carbonique libéré étant déterminés à partir de cette différence de pression, et on répète le traitement thermique, comme indiqué par un ralentissement évident de la réaction et par une formation diminuée de gaz, une substance neutre choisie dans le groupe constitué par les polymères hydrosolubles, les alcools supérieurs, les hydrates de acide organique alimentaire et solide et d'un carbonate et/ou d'un bicarbonate d'un métal alcalin ou alcalino-terraux sous vide, dans lequel, pour la passivation de la surface d'au moins un des composants pour l'amener dans un état après un séchage rapide du mélange, autant de fois que nécessaire pour obtenir une passivation de la surtace, carbone et les hydrocolloïdes étant dissoute dans ledit solvant polaire. **5**6. 53 30 35
- Procédé de préparation d'un matériau granulé effervescent contenant au moins un acide organique alinnentaire cristallin et solide et au moins un carbonate d'un métal alcalin ou d'un métal alcalino-terreux produisant du CO_2 par réaction avec ledit acide organique en solution aqueuse, qui comprend les opérations consistant à : 27.
- provoquer une réaction préliminaire d'une portion dudit acide organique et dudit carbonate en solution dans de 'eau et/ou un alcool pour former un produit de réaction prélininaire,
 - talline et procèder à un mélange poussé pour former un premier revêtement par réaction avec lesdits cristaux ajouter ledit produit de réaction préliminaire à une portion additionnelle dudit acide organique sous forme crisd'acide organique et libération d'eau de cristallisation résultante,

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- appliquer au moins un revétement additionnel comprenant ledit carbonate sur les cristaux d'acide organique avec ledit premier revêtement adhérant à ceux-ci; et
- terminer la réaction après que le dernier revêtement a été appliqué, par un séchage, une substance neutre chaisie dans le groupe constitué par les polymères hydrosolubles, les alcools supérieurs, les hydrates de carsone et les hydrocolloides étant ajoutée audit produit de réaction préliminaire.